

UCLA

UCLA Previously Published Works

Title

A Parietal Biomarker for ADHD Liability: As Predicted by the Distributed Effects Perspective Model of ADHD

Permalink

<https://escholarship.org/uc/item/3674q031>

Journal

Frontiers in Psychiatry, 6(MAY)

ISSN

1664-0640

Authors

Hale, T Sigi
Wiley, Joshua F
Smalley, Susan L
[et al.](#)

Publication Date

2015

DOI

10.3389/fpsyt.2015.00063

Peer reviewed

A parietal biomarker for ADHD liability: as predicted by the distributed effects perspective model of ADHD

T. Sigi Hale^{1*}, Joshua F. Wiley², Susan L. Smalley¹, Kelly L. Tung¹, Olivia Kaminsky¹, James J. McGough¹, Ashwin M. Jaini¹ and Sandra K. Loo¹

¹ Department of Psychiatry and Biobehavioral Sciences, University of California Los Angeles Semel Institute for Neuroscience and Human Behavior, Los Angeles, CA, USA, ² Department of Psychology, University of California Los Angeles, Los Angeles, CA, USA

OPEN ACCESS

Edited by:

Venkata Bharadwaj Kollu,
Creighton University, USA

Reviewed by:

Karen M. Smith,
University of Louisiana at Lafayette,
USA

Katerina Maniadaki,
Technological Educational Institute of
Athens, Greece

*Correspondence:

T. Sigi Hale,
University of California Los Angeles
Semel Institute, 760 Westwood Plaza,
Room 48-228B, Los Angeles,
CA 90024, USA
sig@ucla.edu

Specialty section:

This article was submitted to Child
and Neurodevelopmental Psychiatry,
a section of the journal
Frontiers in Psychiatry

Received: 27 February 2015

Accepted: 11 April 2015

Published: 07 May 2015

Citation:

Hale TS, Wiley JF, Smalley SL,
Tung KL, Kaminsky O, McGough JJ,
Jaini AM and Loo SK (2015) A parietal
biomarker for ADHD liability: as
predicted by the distributed effects
perspective model of ADHD.
Front. Psychiatry 6:63.
doi: 10.3389/fpsy.2015.00063

Background: We previously hypothesized that poor task-directed sensory information processing should be indexed by increased weighting of right hemisphere (RH) biased attention and visuo-perceptual brain functions during task operations and have demonstrated this phenotype in ADHD across multiple studies, using multiple methodologies. However, in our recent distributed effects model of ADHD, we surmised that this phenotype is not ADHD specific, but rather more broadly reflective of any circumstance that disrupts the induction and maintenance of an emergent task-directed neural architecture. Under this view, increased weighting of RH-biased attention and visuo-perceptual brain functions is expected to generally index neurocognitive sets that are not optimized for task-directed thought and action, and when durable expressed, liability for ADHD.

Method: The current study tested this view by examining whether previously identified rightward parietal EEG asymmetry in ADHD was associated with common ADHD characteristics and comorbidities [i.e., ADHD risk factors (RFs)].

Results: Barring one exception (non-right handedness), we found that it was. Rightward parietal asymmetry (RPA) was associated with carrying the DRD4-7R risk allele, being male, having mood disorder, and having anxiety disorder. However, differences in the specific expression of RPA were observed, which are discussed in relation to possible unique mechanisms underlying ADHD liability in different ADHD RFs.

Conclusion: Rightward parietal asymmetry appears to be a durable feature of ADHD liability, as predicted by the Distributed Effects Perspective Model of ADHD. Moreover, variability in the expression of this phenotype may shed light on different sources of ADHD liability.

Keywords: attention, ADHD, laterality, asymmetry, parietal, risk factors, liability, DRD4

Introduction

We recently presented a distributed effects perspective (DEP) model of ADHD that addresses specific challenges associated with the elucidation and treatment of dimensionally defined psychiatric disorders (title: A Distributed Effects Perspective of Dimensionally Defined Psychiatric

Disorders: And Convergent versus Core Deficit Effects in ADHD) (1). The model presents a novel approach to conceptualizing brain–function pathology in dimensionally defined disorders (e.g., distributed versus convergent deficit effects) and makes several predictions regarding abnormal brain function in ADHD. The current study tests one of its key predictions. Below, we present a brief summary description of: (1) the background conceptual framework, (2) research underlying the models genesis, (3) a cursory description of the model, and (4) the rationale for the current empiric study. The overarching goal of this work is to test and refine the DEP model of ADHD, and as such, we present and discuss this study in that context.

The Conceptual Framework

Right hemisphere (RH) specialization is well established for top–down task-directed attention functions, such as vigilance, sustained, and selective attention (2–7). It is also reported for bottom–up functions, such as: detection of sequence breaking novel objects (8), automatic assessment of object relevance (9), automatic perceptual-integrative category learning (10, 11), esthetic analysis (12), and within-category feature discrimination (13). Within these top–down and bottom–up domains, we can further distinguish RH processing that supports fast identifications versus in-depth sensory analysis of stimuli. For instance, top–down processing can facilitate quick stimulus identification using the minimal sensory detail required (i.e., effortful categorizations), or the effortful scrutiny of stimulus details (14). Likewise, bottom–up processing can facilitate fast automatic detection of behaviorally relevant content in our surroundings (i.e., automatic categorizations), or fluid/unguided sensory-immersive experience (15, 16). In short, multiple forms of RH contribution to visual processing can be distinguished; in our work, we have emphasized four specific aspects: (1) top–down task-directed categorizations, (2) top–down task-directed scrutiny of details, (3) bottom–up automatic categorizations, and (4) bottom–up sensory-immersive.

The current line of research began with the precept that complex task-directed actions heavily rely on the first of these (i.e., making fast top–down categorizations of stimuli, using the minimal sensory detail required). We conceptualized this as “task-specialized” sensory information processing, and considered it to reflect variable mixtures of selective, sustained, and/or vigilance related functions depending on the nature of a given task. We hypothesized that ADHD involves a specific deficit for, and/or reduced expression of this task-specialized manner of sensory processing, with a proportional increased expression of non-task-specialized forms (items 2–4 above). We posited that this circumstance should result in mixed expression of: (a) unneeded scrutiny of visual details, (b) increased attentional shifting to off-task content, and/or (c) task-inappropriate orientation toward sensory-immersive processing. The net effect being an increased attentional effort and exposure to visual sensory content beyond what is strictly required to perform task operations. We conceptualized this as “visual sensory overflow” (in relation to task objectives), and predicted that it would be indexed by a relative increased weighting of visuo-perceptual versus verbal-categorical processing during task challenges.

Supporting Literature

We have performed and published three behavioral laterality (17–19), two fMRI (20, 21), and four EEG (22–25), studies examining this view. All produced convergent findings in support of increased RH contribution to sensory processing in ADHD. The behavioral laterality studies showed greater RH contribution to processing task-stimuli, associated left hemisphere (LH) linguistic deficits, and abnormal interhemispheric interaction (17–19). This work also suggested that RH-biased processing in ADHD reflects abnormal brain-state orientation (rather than capacity limitations) (19), bears advantages for RH specialized signal detections (negative tone of voice) (19), impacts high-order cognition (17), and is linked to a superior capacity to inhibit prepotent LH responsiveness (19). The fMRI and EEG studies demonstrated that RH bias in ADHD is mainly evident during sub-executive operations (20), exhibits stronger expression with greater ADHD family loading (23), and is identified in visual (21), inferior and temporal-parietal (22, 25), and frontal (24) brain regions. Moreover, a robust and literature-consistent (26, 27) biomarker was identified. ADHD subjects exhibited pronounced rightward EEG beta asymmetry (16–21 Hz) in inferior parietal brain regions during the Conner’s continuous performance test (CPT) (25). We have replicated this finding (22), and found it to also be present in temporal-parietal regions, along with greater unilateral right-sided beta power in the superior parietal lobe (SPL) (22). This study also replicated an effect showing abnormally weak association between inferior and temporal-parietal beta asymmetry in ADHD. Finally, our most recent fMRI study identified rightward biased visual-cortical function in ADHD, and showed that this characteristic is linked to multiple distributed network systems, including the default mode and the right-lateralized ventral attention networks (21).

Although not yet widely understood, the above findings are well aligned with extant ADHD literature. Abnormal information processing in ADHD is now well established (28, 29), with slow naming speed identified (30–37). Previous behavioral laterality studies have indicated greater RH contribution (38, 39). Functional imaging studies at rest or during simple (i.e., sub-executive) challenges have shown a pattern of reduced LH (40–43) and/or increased RH contribution (20, 24, 44–46). Recent diffusion tensor imaging studies have reported greater RH parietal (47) and frontal (48) fractional anisotropy. A lack of normally occurring $L > R$ asymmetry in prefrontal cortical convolution complexity has also been reported (49), as well as increased RH visual cortex volumes (50). Finally, abnormal posterior corpus callosum size (51) and function (45, 52–54), including abnormally fast left-to-right callosal transfer times (52), clearly implicate some form of abnormal integration of verbal and non-verbal sensory operations in ADHD.

With complex executive function (EF) tasks the literature is more variable, showing diffuse effects mainly consistent with variable weaknesses across multiple brain systems (55–58). Nevertheless, several studies have found an increased association between ADHD behavioral performance and right-sided brain structure and/or function (59–66), and EEG studies that directly examined activation asymmetry and/or left-right hemisphere differences have consistently shown a $R > L$ pattern in posterior brain regions (23–25, 26, 27, 44, 45, 67). A recent meta-analysis of ADHD

functional imaging also reported hyper-activation of the strongly right-lateralized ventral attention network (VAN), noting it may be related to increased bottom-up visuo-perceptual processing of task-extraneous stimuli (28). In this vein, Fassbender and Schweitzer (68), in an earlier review of ADHD brain imaging literature, suggested that ADHD involves an increased reliance on neuroanatomy associated with visual/spatial and motoric (versus verbal) processing.

Finally, increased RH relative to LH contribution may be consistent with common ADHD characteristics, such as: increased leftward motor preference (69, 70), which has been linked to greater RH contribution to language (71, 72); being male (73), as males tend to exhibit better visual-spatial and poorer linguistic abilities compared to females (74, 75), and dysregulated dopamine and noradrenergic function (76), as these systems appear to exhibit a degree of LH and RH specialization respectively (77).

In sum, abnormal information processing is now well established in ADHD. Our work, in association with the above literature, strongly suggests that this abnormality is linked to an atypical increased weighting of RH visuo-perceptual versus LH verbal-categorical processing.

The DEP Model of ADHD

The distributed effects perspective (DEP) model of ADHD asserts that task-directed brain functioning, including task-specialized sensory information processing, is facilitated by the induction and maintenance of an emergent neural architecture (neurocognitive network or brain state) that optimizes multiple distributed systems toward task-directed thought and action (78). The model refers to this emergent neural architecture as a task-directed adaptive processing state (TD-APS) and conceptualizes it to be comprised of four primary computational nodes: (1) verbal working memory (VWM) to sequence, direct, maintain, and update task directives (with possible support from spatial working memory to model integrated plan steps) (79–82), (2) spatial working memory (SWM) to generate predictive sensory models to help bias downstream processing toward task stimuli (83, 84), (3) fast perceptual identification of task-relevant content (14, 16), and (4) translation of perceptual content into verbal articulatory codes that can be readily integrated with, and used to update, task-plans in VWM (85). This system covers four essential task operations: planning, sensory modeling, perceptual encoding, and verbal encoding. It is conceived to have the primary internal objective of orchestrating task directives, and the primary external objective of asserting top-down task-directed control over sensory information processing. The induction and maintenance of this system is expected to coincide with the suppression of automatic sensory and motoric responsivity (86), and requires the dynamic coordination of multiple distributed network systems (e.g., default mode, dorsal and ventral attention, etc.) [For full model description, see Ref. (1)].

The DEP model suggests that ADHD symptom sequelae manifest from TD-APS system impairment, no matter the cause, and that ADHD heterogeneity reflects the diversity of ways in which any such complex distributed system may come to be compromised. Under this view, common ADHD characteristics and/or comorbidities are predicted to reflect common factors that imperil TD-APS system functioning (e.g., sleep-disorder, novelty seeking

temperament, day-dreaming temperament, anxiety, mood, linguistic impairment, working memory impairment, slow processing speed, etc.). Furthermore, “pure ADHD” is conceived to reflect incidences where TD-APS system impairment is both life-disruptive and linked to subclinical causal impairments to its constituent elements. If such impairments are clinically manifest (e.g., impaired verbal encoding), an alternative primary and/or comorbid diagnosis is expected (e.g., dyslexia, slow processing speed, etc.).

An additional important assertion of the model is that variable causal deficits to the TD-APS system are expected to produce “convergent deficit effects” that reflect the system being less able to achieve its primary computational goals. Specifically, TD-APS system impairment, no matter the cause, is expected to result in: (1) a reduced capacity to orchestrate task directives (internal-convergent deficit), indexed by poor working memory and EF-control; (2) a reduced capacity to assert top-down task-directed control over sensory information processing (external-convergent deficit), indexed by increased weighting of RH-biased visuo-perceptual versus left hemisphere (LH) biased verbal-categorical processing; and (3) a general loss of TD-APS system stability (general-convergent deficit), indexed by greater performance variability.

Each of these deficit effects has been established in ADHD [for review, see Ref. (1, 87–91)]. However, the DEP model asserts that they are not ADHD specific, but rather that they are general indicators of poor TD-APS system functioning. If true, the proposed convergent outcomes should be generally present with any circumstance that is tied to poor task-directed brain functioning. Moreover, when such circumstances are time-extended and/or trait-link, they should impart risk for ADHD and be more frequently present in ADHD samples (i.e., common ADHD characteristics and comorbidities).

The Current Study

The current study tests the assertion that previously identified RH-biased sensory information processing in ADHD is reflective of the DEP model’s proposed external-convergent deficit effect. If true, this characteristic (RH-biased processing) should be directly linked to common ADHD characteristics and comorbidities [i.e., ADHD risk factors (RFs)]. To test this hypothesis, we used a large previously collected sample of ADHD children to examine the impact of: (1) being male (73), (2) carrying at least one copy of the DRD4 7 repeat allele (92), (3) having comorbid mood disorder (93), (4) having comorbid anxiety disorder (93), and/or (5) having reduced right-sided motor dominance (69, 70) on rightward parietal EEG asymmetry, and whether this phenotype is associated with ADHD symptoms. As a secondary behavioral validation of RH-biased brain function, we also tested lateralized signal detection using a dichotic listening paradigm. This set of ADHD RFs reflects those that were identified in the literature to be associated with ADHD and that were present in sufficient numbers in our sample to justify analysis.

As a secondary objective, we also performed exploratory analyses to test whether individual ADHD RFs might be associated with different underlying causal impairments to the proposed TD-APS system. We reasoned that if ADHD RFs are associated with unique

causal deficits to this system (distributed deficit effects), rightward parietal EEG asymmetry (the convergent outcome) should exhibit different patterns of association to metrics that tax TD-APS system constituent elements. For example, if the primary mechanism underlying TD-APS system weakness among males is reduced verbal ability relative to females, then male-related increases in rightward parietal EEG asymmetry should exhibit an association to verbal metrics. To explore this idea, for each ADHD RF, we examined the pattern of association between rightward parietal EEG asymmetry and a battery of temperament and cognitive metrics. We also conceptualized that different “points of deficit origination” across TD-APS brain-system constituent elements could result in variant expression of RH-biased brain function, as indexed by variability in the frequency, location, and testing-condition of rightward parietal EEG asymmetry in ADHD RFs.

Materials and Methods

Subjects

The sample consisted of 274 children from an earlier ADHD family genetics study (94, 95). Subjects came from 169 families and comprised of: 79 singletons, 77 sib-pairs, 11 trios, and two quads. After receiving verbal and written explanations of study requirements, a parent and the participating child provided written informed consent/assent, as approved by the UCLA Institutional Review Board. To screen for ADHD and other psychiatric disorders participating children and their mothers were interviewed using the semi-structured interview of the Kiddie Schedule for Affective Disorder and Schizophrenia for School-Age Children-Present and Lifetime Version (K-SADS-PL) (96). Autism was ruled out via the Social Communication Questionnaire (97). All interviews were conducted by clinical psychologists or highly trained interviewers with extensive experience in psychiatric diagnoses. “Best estimate” diagnoses were determined after individual review of diagnoses, symptoms, and impairment level by a board certified child psychiatrist (98). Inter-rater reliabilities were computed with a mean weighted kappa of 0.84 across all diagnoses with a greater than 5% occurrence in the sample. Subjects were excluded based on the following criteria: currently taking psychoactive medication, past or current documented neurological disorder, a significant head injury resulting in concussion, a diagnosis of schizophrenia or autism, or an estimated Full Scale IQ < 80. Inclusion criteria for the present study required a current diagnosis of ADHD. Subjects on stimulant medication were asked to discontinue use for 24 h prior to their visit.

ADHD RFs

Identification of subjects with comorbid mood was based on a definite diagnosis of at least one current mood disorder as assessed by direct interview using the Schedule for Affective Disorders and Schizophrenia-Lifetime Version (SADS-LAR). Identification of subjects with comorbid anxiety required definite diagnosis of at least two current anxiety symptoms as assessed by direct interview using SADS-LAR.

A DRD4 7R group was identified based on subjects possessing at least one copy of the seven-repeat allele (ref). Blood samples were collected from subjects, and DNA was isolated using

the Puregene Kit following the manufacturer’s recommendations (Gentra Systems, Minneapolis, MN, USA). DRD4 polymorphisms (48-bp VNTR and 120-bp repeat) were genotyped according to standard procedures and have been reported elsewhere (99). Since the DRD4-7-repeat allele is thought to be the “risk” allele for ADHD, ADHD subjects were stratified according to whether they possessed at least one copy of the seven-repeat allele (7R) or not (not 7).

Handedness was assessed with a shortened version of the Edinburgh Handedness Inventory (100). This scale uses seven questions regarding hand preference and produces scores ranging from negative 14 (indicating maximum left-handedness), to positive 14 (indicating maximum right handedness). Due to insufficient sample size, we could not produce a purely left-handed RF group. Instead, scores from the Edinburgh Handedness questionnaire were dichotomized so that scores ranging from 8 to 14 indicated “strong right handedness,” and scores less than 8 indicating “not definite right handedness.”

Using a score of less than two standard deviations below the mean on any or our standardized measures of reading, phonologic, or spelling ability produced a small sample ($n = 16$) comprised of variable linguistic weaknesses. Hence, we chose to not include reading disability as an ADHD associated RF in this study. There were no significant differences in linguistic ability based on RF status, as assessed by measures of reading, spelling, phonological, and vocabulary skill (see **Table 2** for description of linguistic measures). Subject demographic information is presented in **Table 1**.

General Testing Procedures

Typical testing procedures for the UCLA ADHD genetics study involved a mother and/or father and two ADHD affected offspring coming to UCLA for a single visit (although fathers were often tested on a separate day). During the visit, each family member underwent a clinical, cognitive, and EEG testing battery, with the order of delivery of each component determined by logistical considerations. However, during EEG testing the Conners’ CPT (101) was delivered first, followed by additional conditions that are not reported. Recordings were performed in a small, private room with a sole male technician administering the protocol.

TABLE 1 | Sample demographics ($n = 274$).

Measures	Statistic
Age	$\bar{x} = 10.9$, std = 3.4
Estimated IQ	$\bar{x} = 111.3$, std = 15.5
ADHD Type	152 C, 113 I, 9 H
ADHD risk factors	
DRD4 7R	88 with 7R allele
Males	177 Males
Mood	26 Affected
Anxiety	43 Affected
Not strong right handed	31

Estimated full IQ: estimated from block design and vocabulary subtest of WAIS-R; ADHD type: C, combined; I, inattentive; H, hyperactive; Mood reflects definite diagnosis of at least 1 current mood disorder as assessed by direct interview using SADS-LAR (see text for reference). Anxiety reflects definite diagnosis of at least 2 current anxiety symptoms as assessed by direct interview using SADS-LAR (see text for reference).

TABLE 2 | Summary of behavioral measures.

Instrument	Measures	Abbrev.	Reference
ADHD symptoms			
Schedule for affective disorder and schizophrenia for school-age children-present and lifetime version (KSADS-PL)	Hyperactive/inattentive symptom counts	KSAD-H KSAD-I	(96)
Swanson, Nolan, and Pelham-IV (SNAP-IV) parent rating scale	4 point Likert scale assessment of hyperactivity and inattention	SNAP-H SNAP-I	(148)
Temperament			
Child behavior checklist (CBCL) social problem scale, plus two recently identified factor items.	Social problems Social immaturity ^a Peer rejection ^a	Soc Prob Soc Imm Peer Rej	(104, 105)
The junior temperament and character inventory scale (JTIC)	Novelty seeking Harm avoidance Reward dependence Persistence Self-directedness Cooperativeness Self-transcendence	NS HA RD PR SD CP ST	(136)
Cognition			
Wechsler intelligence scale for children, third edition (WISC-II)	Block design Vocabulary Coding Arithmetic Digit span forward max Digit span forward acc Digit span backward max Digit span backward acc Spatial span forward max Spatial span forward acc Spatial span backward max Spatial span backward acc	Blocks Vocab Coding Arith DSF-max DSF-acc DSB-max DSB-acc SSF-max SSF-acc SSB-max SSB-acc	(149)
Peabody individual achievement test-revised (PIAT-R)	Reading recognition Spelling	Read_rec Spell	(150)
Woodcock Johnson-revised, word attack subtest (WJ-R)	Phonologic processing	Phono	(151)
Stroop task	Color naming Word naming Interference	St-Color St-Word St-Inter	(152, 153)
The trail making test – A and B	Trails A Trails B	Trails A Trails B	(154)
Conner's continuous performance test II (CPT-II)	Commissions Omissions Hit reaction time Hit RT standard error Sensitivity Bias	Commiss Omiss Hit RT Hit RTSE D-prime Beta	(101)
Spatial working memory (SWM)	Load 1 accuracy Load 1 reaction time Load 1 RT standard dev. (+ loads: 3, 5, 7)	SWM L1 Acc SWM L1 RT SWM L1 RTSD (+ load: 3, 5, 7)	(107)
Dichotic listening emotion/words	Left-ear words (Acc/RT) Right-ear words (Acc/RT) Left-ear emotion (Acc/RT) Right-ear emotion (Acc/RT)	LE-word RE-word LE-emot RE-emot	(127)

^a Novel indices from CBCL derived from factor analysis (see text for reference).

EEG Procedures

EEG recording was carried out using 40 silver chloride electrodes using the International 10/20 locations and was referenced to an average of singles recorded separately at each ear lobe. Eye movements were monitored by electrodes placed on the outer canthus of each eye for horizontal movements and by electrodes above the eye for vertical eye movements. EEG recording consisted of 2 baseline conditions lasting 5 min each [eyes open (EO) and eyes

closed (EC)] and a cognitive activation condition lasting 15 min (Conners CPT-II) (102).

Continuous EEG data were subjected to automatic artifact detection via MANSKAN software (SAM Technology, Inc., San Francisco, CA, USA) designed to identify dead and bad channels, vertical and horizontal eye movements, saturation, muscle and movement artifact, and line frequency noise. Subsequent to this automated procedure an experienced EEG technician then

visually inspected all data and identified any residual contaminants. Next, continuous EEG was broken into 1-s epochs and artifact-containing epochs were removed on a channel specific basis. Remaining artifact free epochs were then Fast Fourier Transformed (FFT) using MANSKAN EEG software, which uses a Welch's Periodogram approach (103). We specified 1-s data segments, with 50% overlap, and a Hanning Windowing function to generate spectral content at a 1 Hz resolution. Spectral data were then averaged for each condition (EC, EO, CPT), and EEG power (mv^2) from 1 to 21 Hz was exported in 1 Hz bins (e.g., 0–1, 1–2, . . . 20–21). Technicians involved in the EEG recording and processing were blind to ADHD diagnostic status.

For the current study, we evaluated the EEG frequency bands that were previously associated with abnormal EEG asymmetry in ADHD (23–25, 26, 27, 44, 67). Absolute power between 8–10 Hz, 10–12 Hz, 12–16 Hz, and 16–21 Hz frequencies were averaged for each electrode composing “low” and “high” alpha and beta measures, respectively (A1, A2, B1, B2). Our primary interest was to evaluate relative $R > L$ parietal brain activation in RF groups. We therefore utilized measures of power asymmetry rather than examining group differences separately in each hemisphere. Laterality indices (LIs) were generated for three homologous right–left parietal electrode pairs (TP8-TP7, P4-P3, P8-P7) using the following standard calculation: $[(R - L)/(R + L) * 1000]$. Greater values for LIs mean greater rightward asymmetry. EEG asymmetry indices are notated as follows: condition (EC, EO, CPT), frequency (A2, A2, B1, B2), parietal region (TP8-7, P4-3, P8-7), for example, EC A1 TP8-7.

Clinical and Behavioral Measures

Commonly used measures are listed in **Table 2**. Selected measures are also described here. Note: some tasks were added at different stages of the ADHD family genetics study, which is reflected by smaller sample sizes for some measures (e.g., CPT task).

Temperament

Social functioning characteristics were assessed using the mother's report on the Social Problem scale of the child behavior checklist (CBCL) (104), as well as two previously identified factors from this scale (105). The first factor called “social immaturity” included items 1 (acts young), 11 (clings), 62 (clumsy), and 64 (prefer young). The second factor called “peer rejection” included items 25 (not get along), 48 (not liked), and 39 (teased). The Junior Temperament and Character Inventory scale (JTCI) was also used. This measures individual differences on seven personality dimensions: four dimensions of temperament (novelty seeking, harm avoidance, reward dependence, persistence), and three dimensions of character (self-directedness, cooperativeness, self-transcendence).

Conner's Continuous Performance Test II

The CPT requires subjects to monitor a central fixation on a computer screen while single capital letters are sequentially and centrally presented during six continuous blocks of 20 trials with either 1, 2, or 4 s inter-stimulus intervals (ISIs) (two blocks for each ISI). The order of ISI-block presentation is randomized within subjects. The task requires subjects to press the space bar using

their dominant hand with every letter presentation except for the letter “X.” The “X” occurs on 10% of the trials within a given ISI block. Behavioral performance was assessed using the following standard measures (106): (1) commission errors: a failure to inhibit response when an “x” is presented, (2) omission errors: a failure to respond when any letter other than “x” is presented, (3) hit reaction time: response time for all letters other than “x,” (4) hit reaction time standard error: reaction time variability, (5) response bias: signal detection measure (beta) indicating impulsive versus conservative response styles, and (6) sensitivity: signal detection measure (d') indicating accuracy adjusted for false alarms.

Spatial Working Memory

In this spatial delay response task (SDRT) (107), subjects were shown a target array of 1, 3, 5, or 7 yellow circles positioned pseudo-randomly around a fixation point. Then, after a fixed delay, they were shown a single green circle and had to indicate if it was in the same position as any of the previous yellow circles. Trial events included a 500-ms initial period of blank screen, 500 ms of a fixation point, a 2-s target array presentation, a 3-s delay period (with fixation), and a 3-s fixed response interval. Fifty percent of the trials were true positive, and 50% were true negative. Responses (yes or no) and reaction times were recorded for each trial. The number of correct responses, mean reaction time, and reaction time variability were recorded.

Dichotic Listening for Emotions and Words

This task requires subjects to detect an emotional tone of voice or to identify a word during dichotic presentations of four words (bower, dower, power, tower) spoken in four different emotional tones (happy, sad, angry, neutral) by a male speaker. Stimuli for the “word” and “emotion” tasks are identical and only the instructions to the subject change between conditions. Because dichotic presentations suppress ipsilateral auditory channels, the stimuli in each ear project to the opposite hemisphere via fibers that cross at the superior olive and inferior colliculus (108, 109). Thus, if each hemisphere can process the stimuli presented in the contralateral ear [direct access: (110)], then subjects' ability to detect signal in the right ear, for instance, is indicative of left hemisphere (LH) competence. Based on previous work, we used nominated targets “bower” and “sad,” as these show the most robust LH and RH specialization, respectively. Moreover, with these stimuli, right-ear presentations of the “sad tone of voice” (projecting to the LH) are expected to require a left-to-right callosal transfer to the RH (111). Likewise, left-ear presentations of the word “bower” (projecting to the RH) are expected to involve right-to-left transfer to the LH, although some RH competence for the word signal is also expected (111). This task consistently demonstrates a left-ear (RH) advantage for processing emotional intonation and a right-ear (LH) advantage for processing words (112).

Data Analytic Procedures and Results Presentation Overview

We first present analyses to characterize ADHD RF groups with respect to clinical, temperament, and cognitive characteristic. Next, analyses aimed at testing the DEP model of ADHD are

presented and discussed in two parts. Part 1 addresses our primary aim of examining whether RH bias is a general feature of ADHD RFs. Part 2 addresses our secondary/exploratory aim of examining whether ADHD RFs show different patterns of association between RH bias and temperament and/or cognitive characteristics. Section specific discussions are presented for Parts 1 and 2, followed by a general discussion. A summary of all study findings is provided in the supplement (Table S3 in Supplementary Material).

Non-Independence

Non-independence in our data (sibling pairs) is distributed across the levels of our grouping variables, making it somewhat ambiguous with respect to increasing or decreasing our ability to detect RF group differences. For instance, with regard to the “being male” RF, siblings are just as likely to be the same or different genders. If different, non-independence hurts our capacity to detect a gender difference – if the same, it helps. Moreover, because our emphasis was to uncover conceptually meaningful patterns, rather than individual effects, our interpretation of results is conceptually insulated against type 1 error. Nevertheless, steps were taken to reduce this risk (described below).

Multiple Testing

In primary analyses of rightward parietal asymmetry (RPA) in ADHD RFs (Part 1), the multivariate multilevel modeling approach limited the number of tests and accounted for non-independence (by modeling family membership). Analyses to characterize our samples, and secondary/exploratory analyses (Part 2), utilized large batteries of cognitive and temperament metrics (41 measures – see **Table 3**), and so were vulnerable to type 1 error. However, as noted, our goal in performing these analyses was to examine patterns of results among different RFs (i.e., we do not interpret individual findings). Nevertheless, to help guard against false positives diluting, distorting, or disrupting otherwise coherent patterns, and to reduce the complexity of this large set of findings, results for temperament and cognitive outcomes are restricted to a *p* threshold of 0.015. Symptom metrics included only four measures, and because all subjects had ADHD, we expected limited variability in these measures. Hence, we did not restrict symptom findings to a 0.015 threshold.

Results

Sample Characteristics

Univariate ANOVA (SPSS v21) was used to examine each ADHD RF with respect to: ADHD symptoms, temperament, and cognitive abilities. For these analyses, a clinical, temperament, or cognitive measure of interest was entered as the dependent variable, an ADHD RF of interest was entered as the grouping variable, with age plus the remaining ADHD RFs entered as covariates in order to assess the unique impact of the ADHD RF being tested.

ADHD Symptoms

DRD4 7R carriers exhibited fewer inattentive symptoms. Male, mood, and anxiety RFs were associated with greater hyperactivity. Non-right handedness had no effect on symptoms (**Tables 3** and **Summary Table 4**).

TABLE 3 | General characteristics by ADHD risk factor type.

RFs and measures	No RF		With RF		Statistics			With RF
	\bar{x}	se	\bar{x}	se	df	<i>f</i>	<i>P</i> -value	
Symptoms (handedness no effects)								
<i>DRD4</i>								
KSAD-I	8.1	(0.13)	7.5	(0.19)	1,104	6.8	0.010	Less
<i>Sex</i>								
KSAD-H	5.0	(0.26)	5.7	(0.21)	1,260	5.2	0.024	More
SNAP-H	1.53	(0.07)	1.76	(0.06)	1,255	6.4	0.012	More
<i>Mood</i>								
KSAD-H	5.3	(0.17)	6.7	(0.57)	1,260	5.6	0.019	More
<i>Anxiety</i>								
KSAD-H	5.3	(0.18)	6.2	(0.43)	1,260	3.8	0.053	More
Temperament (<i>P</i> threshold: 0.015: DRD4 7R, handedness no effects)								
<i>Sex</i>								
RD	5.3	(0.21)	4.4	(0.17)	1,216	10.0	0.002	Less
CP	15.9	(0.35)	14.2	(0.28)	1,216	14.9	0.0001	Less
ST	5.8	(0.23)	5.0	(0.18)	1,216	7.2	0.008	Less
NS	7.6	(0.38)	9.4	(0.30)	1,216	13.2	0.0003	More
<i>Mood</i>								
ST	5.5	(0.15)	4.0	(0.51)	1,216	7.6	0.006	Less
Peer Rej	1.0	(0.09)	1.9	(0.31)	1,237	7.5	0.007	More
<i>Anxiety</i>								
HA	7.5	(0.31)	10.8	(0.74)	1,216	16.0	0.00009	More
ST	5.2	(0.15)	6.2	(0.36)	1,216	6.1	0.014	More
Soc Imm	1.7	(0.12)	2.8	(0.30)	1,235	11.0	0.001	More
Soc Prob	2.7	(0.19)	4.4	(0.48)	1,233	10.4	0.001	More
Cognition (<i>P</i> threshold: 0.015: handedness no effects)								
<i>DRD4</i>								
Info	12.3	(0.24)	10.9	(0.35)	1,262	11.4	0.001	Worse
SSF-max	4.9	(0.08)	5.4	(0.12)	1,255	9.7	0.002	Better
<i>Sex</i>								
CPT D-prime	0.39	(0.03)	0.26	(0.02)	1,118	15.7	0.0001	Worse
Coding	10.5	(0.31)	9.0	(0.25)	1,261	14.3	0.0002	Worse
<i>Mood</i>								
CPT Commiss	17.6	(0.90)	21.6	(0.70)	1,118	12.1	0.001	Worse
<i>Anxiety</i>								
CPT Commiss	19.5	(0.61)	23.1	(1.3)	1,118	6.2	0.014	Worse
CPT D-prime	0.33	(0.02)	0.22	(0.04)	1,118	6.7	0.011	Worse
Info	12.1	(0.21)	10.7	(0.50)	1,262	6.2	0.013	Worse

Univariate analysis of variance was used to assess the impact of ADHD RFs on ADHD symptoms, temperament, and cognition. Each analysis reflects the effects of a specific RF after adjusting (i.e., co-varying) for the effects of age and the remaining additional RFs. See dependent measures list (**Table 2**) for description of measures.

Temperament

DRD4 7R carriers and non-right handers showed no effects. Being male was associated with a less cooperative and more novelty seeking characteristics. Mood and anxiety sufferers exhibited social functioning liabilities. Anxiety also showed a strong association with harm avoidance (**Table 3** and **Summary Table 4**).

Cognitive Abilities

Non-right-handers showed no effects. DRD4-7R carriers showed an enhanced ability for SWM, and worse “general knowledge,” as indexed by the WISC III information task. Males,

TABLE 4 | Summary of ADHD risk factor effects on clinical and cognitive measures.

Assessments	DRDR 7R	Males	Mood affected	Anxiety affected	Non-right handed
Symptoms	Less IA	More H	More H	More H	None
Temperament	None	More NS Less RD Less CP Less ST	More Peer Rej Less ST	More HA More ST More Soc Imm More Soc Prob	None
Cognition	Worse Info Better SSF-max	Worse D-prime Worse coding	Worse Commiss	Worse Info Worse D-prime Worse Commiss	None

IA, inattention; H, hyperactivity.

For description of additional abbreviations, see Table 2.

anxiety, and mood were associated with worse CPT performance. Males showed poor CPT stimulus discrimination. Mood sufferers show poorer CPT stop inhibition. Anxiety sufferers show both poorer stimulus discrimination and stop inhibition, as well as poor “general knowledge” (Table 3 and Summary Table 4).

Part 1: Primary Analyses of Right Hemisphere Bias in ADHD RFs

This section examines whether RH-biased brain function is a general feature of ADHD RFs. Three analyses are performed: (1) examination of parietal EEG asymmetry, (2) examination of RPA association to ADHD symptoms, and (3) examination of dichotic listening behavioral performance.

Step 1: Parietal EEG Asymmetry

We tested whether individual ADHD RFs were associated with increased rightward parietal EEG asymmetry (RPA) using multilevel models (software: R 2.15.2). The first level was the EEG asymmetry measure (i.e., condition and frequency). The second level was the parietal area. The third level was the family. To limit the number of tests we first examined whether there were overall effects of parietal EEG asymmetry (collapsed across specific parietal locations), then significant effects were further decomposed by looking at pairwise comparisons of individual parietal asymmetry indices.

ADHD RFs were shown to impact overall parietal EEG asymmetry. Effects were broadly expressed for DRD4 7R carriers and males, but specific for mood, anxiety, and non-right handedness (Table 5).

Assessment of location-specific asymmetry indices showed that 38 of 39 significant findings exhibited RPA. RPA was broadly expressed in DRD4 7R carriers and males, but specifically expressed in mood and anxiety groups. RPA appeared strongest at the P8-7 LI. Although both DRD4 7R carriers and males showed broad RPA expression, the DRD4 7R group showed asymmetry effects mainly in alpha at the TP8-7 and P8-7 indices, while males showed asymmetry effects mainly in beta (especially beta1) across all three parietal locations. Mood only showed RPA effects during EC in beta2, while anxiety only showed RPA during active visual processing (EO, CPT) in beta1. Individuals who were not strongly right-handed did not show RPA (Table 6 and Summary Table 9).

TABLE 5 | Overall parietal EEG asymmetry by ADHD risk factor type.

Parietal Asymmetry by risk factor	Statistics		
	df	Chi-square	P-value
DRD4 7R			
EC A1	3	36.6	0.0000005
EC A2	3	26.5	0.0000007
EC B1	3	10.34	0.016
EC B2	3	22.6	0.00005
EO A1	3	15.62	0.0013
EO A2	3	11.24	0.010
EO B1	3	10.94	0.012
EO B2	3	11.05	0.011
CPT A1	3	9.40	0.024
CPT A2	3	9.53	0.023
CPT B1	3	7.81	0.050
Sex			
EC A2	3	17.77	0.0005
EC B1	3	13.54	0.003
EC B2	3	9.69	0.021
EO B1	3	18.52	0.0003
CPT A2	3	16.0	0.001
CPT B1	3	16.21	0.001
Mood			
EC B2	3	14.76	0.002
Anxiety			
EO B1	3	10.94	0.012
CPT B1	3	9.44	0.024
Handedness			
EC A2	3	12.51	0.005

Linear multilevel models and tests were to examine the overall effect of RPA by frequency and condition collapsed across three parietal brain regions. The first level was the EEG asymmetry measure (i.e., condition and frequency), the second level was the parietal area (tp8-tp7, p8-p7, p4-p3), and the third level was the family. Then for each RF, we conducted multivariate tests to examine whether it had an overall effect across the three parietal brain regions in the model. See dependent measures list (Table 2) for description of measures.

Step 2: Parietal EEG Asymmetry Association to Symptoms in ADHD RFs

Linear regression (SPSS v21) was used to examine whether RF groups interacted with individual parietal EEG asymmetry indices to predict ADHD symptoms, while controlling for age and other RF groups. Here, a symptom metric was entered as the outcome variable, then age and the “not to be examined” RF groups were entered as predictors, followed by the RF group being examined,

TABLE 6 | Specific parietal EEG asymmetry indices by ADHD risk factor type.

Asymmetry indices	No RF		With RF		Statistics			With RF asym. eff.
	\bar{x}	se	\bar{x}	se	df	t	P-value	
DRD4 7R								Carriers
EC A1 TP8-7	65 (20)		160 (24)		432	3.51	0.0005	R
EC A1 P8-7	70 (21)		232 (26)		432	5.68	0.00000002	R
EC A1 P4-3	50 (18)		96 (21)		432	1.97	0.049	R
EC A2 TP8-7	83 (21)		159 (25)		432	2.76	0.006	R
EC A2 P8-7	94 (22)		230 (27)		432	4.60	0.000006	R
EC B1 P8-7	57 (15)		106 (18)		431	2.47	0.014	R
EC B2 TP8-7	27 (12)		67 (14)		430	2.55	0.011	R
EC B2 P8-7	18 (13)		94 (16)		430	4.26	0.00002	R
EO A1 P8-7	67 (20)		127 (25)		395	2.15	0.032	R
EO A2 TP8-7	51 (21)		109 (26)		395	2.02	0.044	R
EO B1 TP8-7	21 (13)		67 (16)		395	2.58	0.010	R
EO B1 P8-7	33 (14)		70 (17)		395	1.95	0.050	R
EO B2 P8-7	-2 (13)		42 (16)		395	2.46	0.014	R
CPT A1 TP8-7	40 (16)		89 (21)		429	2.10	0.036	R
CPT A1 P8-7	109 (18)		179 (23)		429	2.8	0.005	R
CPT A2 TP8-7	82 (19)		139 (24)		429	2.11	0.035	R
CPT A2 P8-7	152 (21)		234 (27)		429	2.76	0.006	R
CPT B1 TP8-7	0.96 (15)		52 (19)		429	2.50	0.013	R
Sex								Males
EC A2 TP8-7	9 (23)		83 (21)		432	2.90	0.004	R
EC A2 P8-7	-15 (25)		94 (22)		432	3.92	0.0001	R
EC B1 TP8-7	-5 (16)		38 (15)		431	2.34	0.02	R
EC B1 P8-7	-8 (17)		57 (15)		431	3.45	0.0006	R
EC B1 P4-3	7 (12)		42 (11)		431	2.52	0.012	R
EC B2 TP8-7	-5 (14)		27 (12)		430	2.08	0.038	R
EC B2 P8-7	-27 (15)		18 (13)		430	2.65	0.008	R
EO B1 TP8-7	-34 (16)		21 (13)		395	3.23	0.0013	R
EO B1 P8-7	-25 (17)		33 (14)		395	3.18	0.0016	R
EO B1 P4-3	5 (9)		37 (8)		395	3.17	0.0016	R
CPT A2 TP8-7	11 (22)		82 (19)		429	3.23	0.004	R
CPT A2 P8-7	55 (24)		152 (21)		429	3.55	0.0004	R
CPT A2 P4-3	59 (15)		99 (14)		429	2.31	0.020	R
CPT B1 P8-7	9 (16)		72 (14)		429	3.50	0.0005	R
CPT B1 P4-3	18 (9)		48 (8)		429	3.12	0.002	R
Mood								Affected
EC B2 P8-7	18 (13)		112 (27)		430	3.39	0.0007	R
EC B2 P4-3	16 (8)		63 (17)		430	2.75	0.0061	R
Anxiety								Affected
EO B1 TP8-7	21 (13)		67 (23)		395	2.04	0.042	R
EO B1 P8-7	33 (14)		97 (25)		395	2.63	0.009	R
CPT B1 TP8-7	0.96 (15)		58 (27)		429	2.24	0.026	R
Handedness								Weak right-handed
EC A2 TP8-7	83 (21)		-0.99 (41)		432	2.06	0.040	L

Pairwise comparisons of individual parietal asymmetry indices for each ADHD RF, where the above first pass test showed a general effect on parietal asymmetry (see Table 8). Like in the first pass analysis, these pairwise comparisons used linear multilevel models. The first level was the EEG asymmetry measure (i.e., condition and frequency), the second level was the parietal area (tp8-tp7, p8-p7, p4-p3), and the third level was the family. Note: x-bar reflects the model adjusted means. See dependent measures list (Table 2) for description of measures.

the parietal asymmetry index being examined, and their interaction term.

Mood had no effects. All other RFs showed associations between parietal EEG asymmetry and ADHD symptoms, with anxiety showing the strongest and most numerous effects with unique weighting toward the alpha band. The findings generally

demonstrated that RPA was associated with increased symptoms, except for two instances. DRD4-7R carriers showed a link between RPA and fewer inattentive symptoms, and two of eighteen significant effects for anxiety showed a link between RPA and reduced hyperactivity. In males and DRD4-7R carriers, RPA was associated with more hyperactivity. In non-right-handers, it was associated with more inattention. In anxiety, it was associated with more inattention and hyperactivity. In DRD4 7R carriers, males, and non-right handed individuals, a relationship between RPA and ADHD symptoms was mainly present for the P4-3 LI (exclusively so in males) (Table 7 and Summary Table 9).

Step 3: Dichotic Listening Assessment of RH Bias in ADHD RFs

As a secondary behavioral validation of RH-biased processing among ADHD RFs, we examined whether ADHD RFs exhibited RH-biased signal detection during the dichotic listening paradigm, using the same univariate ANOVA procedure described in Part 1.

Mood and non-right handedness showed no effects. All significant effects showed faster responses for RF groups. DRD4 7R carriers were faster for word processing in both ears, but the effect was stronger for left-ear (RH) trials. Moreover, they did not show the standard right-ear (LH) speed advantage for word stimuli, and instead showed a left-ear (RH) advantage. DRD4-7R carriers also showed a speed advantage for detecting right-ear emotion stimuli. This trial type is understood to depend on left-to-right hemisphere transfer of emotion stimuli due to strong RH specialization for these stimuli (111). Males showed a direct left-ear (RH) speed advantage for emotion stimuli, while anxiety sufferers showed the same right-ear speed advantage for emotion stimuli as the DRD4 7R carriers (Table 8 and Summary Table 9).

Part 1: Discussion

We previously hypothesized that non-task-specialized sensory information processing should be indexed by increased weighting of RH-biased attention and visuo-perceptual brain functions during task operations, and demonstrated this phenotype in ADHD across multiple studies, using multiple methodologies [for review, see Ref. (1)]. However, as detailed in the DEP model (1), we surmised that this phenotype is not ADHD specific, but more broadly reflective of any circumstance that disrupts the induction and maintenance of an emergent task-directed neural architecture (i.e., a task-directed adaptive processing state or TD-APS). Under this view, abnormal increased weighting of RH-biased attention and visuo-perceptual brain functions is expected to generally index neurocognitive sets that are not optimized for task-directed thought and action, and when durably expressed, liability for ADHD.

The current study tested this view by examining whether previously identified rightward parietal EEG asymmetry in ADHD (22, 25, 26, 27) is generally associated with common ADHD characteristics and comorbidities (i.e., ADHD RFs). Barring one exception (non-right handedness), we found that it was. RPA was associated with carrying the DRD4-7R risk allele, being male, having mood disorder, and having anxiety disorder. This pattern was additionally supported by behavioral laterality outcomes. However, differences in the specific expression of RPA and behavioral

TABLE 7 | Parietal EEG asymmetry × risk factors predicting ADHD symptoms.

RFs × EEG asymmetry predicts	df	SB	t	P-value	With RF rightward predicts	Continued ...	df	SB	t	P-value	With RF rightward predicts
DRD4 7R						Anxiety continued					
<i>KSAD Inattention</i>						EC A2 P8-7 1,205 0.17 2.1 0.037 More					
EC A1 P4-3	1,214	-0.20	-2.3	0.027	Less	EC B1 TP8-7	1,210	0.16	2.1	0.040	More
EO A1 P4-3	1,200	-0.20	-2.3	0.024	Less	EO A2 P8-7	1,191	0.19	2.3	0.022	More
EO B2 P4-3	1,200	-0.17	-2.0	0.05	Less	EO A2 P4-3	1,198	0.16	2.1	0.040	More
CPT B1 TP8-7	1,195	-0.18	-2.0	0.05	Less	CPT A1 P8-7	1,205	0.17	2.0	0.045	More
<i>KSAD Hyperactive</i>						CPT A2 TP8-7 1,211 0.15 2.0 0.043 More					
EC B1 TP8-7	1,215	0.19	2.3	0.023	More	CPT A2 P8-7	1,205	0.18	2.2	0.030	More
CPT A2 P4-3	1,219	0.22	2.3	0.021	More	CPT A2 P4-3	1,214	0.21	2.6	0.009	More
Male						CPT B1 P8-7 1,205 0.21 2.6 0.010 More					
<i>KSAD Hyperactive</i>						CPT B2 TP8-7 1,211 0.17 2.2 0.026 More					
EC B2 P4-3	1,214	0.20	2.0	0.050	More	CPT B2 P8-7	1,205	0.28	3.9	0.0001	More
<i>SNAP Hyperactive</i>						CPT B2 P4-3 1,214 0.16 2.0 0.047 More					
EC B2 P4-3	1,209	0.28	2.9	0.004	More	<i>KSAD Hyperactive</i>					
CPT A1 P4-3	1,214	0.30	2.7	0.006	More	EO A1 P4-3	1,200	-0.18	-2.3	0.021	Less
Mood						EO A2 P4-3 1,91 -0.34 -3.0 0.003 Less					
<i>Nothing</i>						CPT B1 TP8-7 1,215 0.21 2.80 0.006 More					
Anxiety						Handedness					
<i>KSAD Inattentive</i>						<i>SNAP Inattention</i>					
CPT B2 TP8-7	1,215	0.20	2.8	0.006	More	EC A1 TP8-7	1,210	-0.74	-2.5	0.014	More
<i>SNAP Inattentive</i>						CPT A2 P4-3 1,214 -0.49 -2.1 0.036 More					
EC A1 TP8-7	1,210	0.20	2.4	0.016	More	CPT B1 P4-3	1,214	-0.52	-2.3	0.025	More
EC A1 P8-7	1,205	0.21	2.5	0.013	More	CPT B2 P4-3	1,214	-0.45	-2.5	0.012	More

Linear regression analysis was used to examine the interaction effects of RFs and parietal EEG asymmetry on ADHD symptoms. Each test was adjusted for the effects of age and additional RFs. See dependent measures list (Table 2) for description of measures.

TABLE 8 | Dichotic listening by ADHD risk factor type.

Group and BD measures	No RF		With RF		Statistics			With RF
	\bar{x}	se	\bar{x}	se	df	f	P	
DRD4 7R								
LE-word RT	1065	(19.3)	933	(27.8)	1,167	15.0	0.0001	Faster
RE-word RT	1025	(17.8)	937	(25.6)	1,167	7.8	0.006	Faster
RE-emot RT	1150	(22.9)	1068	(33.0)	1,167	4.0	0.046	Faster
Sex								
LE-emot RT	1136	(22.8)	1072	(20.3)	1,167	4.3	0.040	Faster
Anxiety								
RE-emot RT	1142	(20.4)	1023	(48.6)	1,167	5.0	0.027	Faster

Univariate analysis of variance was used to assess the impact of ADHD RFs on eight dichotic listening measures (left and right performance for word and emotion targets – in accuracy and latency). All significant effects were for latency measures. Each analysis was adjusted for the effects of age and additional RFs. See dependent measures list (Table 2) for description of cognitive measures. Mood and non-right handedness showed no effects.

laterality were observed, which are discussed in Part 2 in relation to possible unique mechanisms underlying ADHD liability in different ADHD RFs.

We view the single outlying handedness result with caution. Due to sample constraints, we did not test left-handedness versus right handedness, but rather the effect of being “strongly” versus “not strongly” right-handed, with the latter group including weakly right-handed, ambidextrous, and left-handed individuals. Moreover, secondary analyses implicated RH-biased processing in the “not strongly right-handed” group. Hence, we think it is

reasonable to refrain from interpreting these conflicting outcomes until a better test of pure left-handedness is performed.

The otherwise clear pattern of RPA across the remaining ADHD RFs demonstrates, as predicted by the DEP model, a connection between RH-biased parietal brain function and liability for ADHD, and by inference, a general reduced capacity for task-directed thought and action. This outcome is well aligned with the established role of right-lateralized parietal brain function in attention and visual sensory information processing.

Although much remains to be elucidated about parietal brain function, multiple lines of research indicate that it plays a key role in processing information in a spatial context (113), with a dorsal-to-ventral distribution of functions related to “vision for action” versus “vision for perception” (114), and a left-to-right distribution of functions related to self-directed motoric and verbal functions (LH), versus the allocation of attention to external sensory content (RH) [for review, see Ref. (115)]. This left-right dimensionality is evident across the superior, inferior, and temporal-parietal regions examined in the current study. For instance, the LH SPL shows specialization for self-initiated fine motor actions such as writing (114, 116), whereas the RH SPL has been associated with spatial orienting (114). Similarly, the LH inferior parietal lobe (IPL) shows specialization for self-initiated grasping of objects (117), while the RH IPL is associated with orienting and sustaining attention toward external stimuli [for review, see Ref. (115, 118)].

Continuing this pattern, the LH temporal-parietal junction (TPJ) shows specialization for verbal articulatory coding (i.e., naming) (85), while the RH TPJ has been associated with stimulus-driven attentional shifting (119, 120). Related to this,

TABLE 9 | Summary of right hemisphere bias assessment in ADHD RFs.

Laterality	DRDR 7R	Males	Mood	Anxiety	Non right-handed
Parietal EEG asymmetry	All rightward 18 Effects Mixed cond. Mostly alpha Mainly TP8-7, P8-7	All rightward 15 Effects Mixed cond. Mostly beta1 Mixed location	All rightward 2 Effects Only EC Only beta2 P4-3 and P8-7	All rightward 3 Effects Only EO, CPT Only beta1 TP8-7, P8-7	1 Leftward EC A2 TP8-7
RPA association to symptoms	Less IA. (4) More H (2)	More H (3)	None	More IA (15) More H (1) Less H (2)	More IA (4)
Dichotic listening validation	RH bias indicated Faster word RT No RE-word RT adv. Faster RE-emot (L-to-R trans. cond.)	RH bias indicated Faster LE-emot	None	RH bias indicated Faster RE-emot (L-to-R trans. cond.)	None

For "RPA association to symptoms", RPA (i.e., greater rightward parietal asymmetry) is associated with the reported outcomes. cond, condition; H, hyperactive; IA, inattentive; effs, effects; RE, right ear; LE, left ear; RT, reaction time; trans., transfer; adv., advantage; emot., emotion; "L-to-R trans. cond." refers to the expectation that emotion stimuli presented in the right ear during this dichotic listening task are understood to undergo callosal transfer from the left to the right hemisphere, which is dominant for processing the emotion stimuli (see task description for details). For description of additional abbreviations, see **Table 2**.

Geng and Vossel (119) have recently argued that the RH TPJ plays a key role in maintaining and updating the neural context by which the relevance of incoming sensory information is vetted. This model implies that greater RH TPJ activation indexes a more flexible attentional and cognitive set (i.e., with more active updating and shifting), while reduced activation indexes a more fixed attentional and cognitive set (i.e., with less active updating and shifting). They also specify that this RH TPJ function likely draws on multiple distributed brain systems and integrates both bottom-up and top-down processing. According to this view, moment-to-moment variability in RH TPJ activation may generally index the relative degree to which individuals are oriented toward a more fixed versus flexible cognitive and attentional set.

We have previously demonstrated that ADHD is associated with abnormal increased rightward asymmetry across each of these parietal regions (P4-P3:SPL, P8-P7:IPL, TP8-TP7:TPJ) (22–25), and importantly, these outcomes could not be attributed to gender, handedness, anxiety, and/or mood. This demonstrates that ADHD involves abnormal increased weighting of right-lateralized parietal brain functions associated with: orienting attention in space (RH SPL), sustaining attention on external sensory content (RH IPL), and possibly having a more flexible attention and cognitive set (RH TPJ). Broadly speaking, these findings demonstrate an increased weighting of external-perceptual versus verbal-motoric parietal brain function in ADHD. They also highlight two possible orthogonal mechanisms that might underlie this characteristic. Greater R > L TPJ activation might index an overly flexible cognitive and perceptual set (a brain state-setting effect), while R > L SPL and IPL activations might be associated with greater attentional effort (possibly tied to compensatory effort and/or the processing of task-extraneous content).

The current study has now demonstrated that this same pattern of R > L parietal brain functions is an independent feature of multiple ADHD RFs. This adds strong support to the view that RPA may generally index: (1) brain function that is not optimized for task-directed thought and action and (2) when durably expressed, liability for ADHD.

Part 2: Secondary Analyses of RPA Association to Temperament and Cognition in ADHD RFs

This secondary/exploratory analysis examined whether ADHD RFs exhibited different patterns of association between RPA and temperament and cognition metrics. It used the same linear regression approach described above, but with temperament and cognition metrics (rather than symptom measures) entered as outcome variables. This analysis helped us to explore whether different mechanisms (i.e., divergent/distributed deficit effects) might underlie RH-biased brain function across different ADHD RFs. General results patterns and summary tables are provided here, with full results details available in the study supplement (Tables S1 and S2 in Supplementary Material).

Parietal Asymmetry Association to Temperament in Different ADHD RFs

RPA showed associations to temperament characteristics in each ADHD RF. Except for males, most RPA associations to temperament occurred in the alpha-frequency band. In males, these effects were mainly in beta. Males, mood, and anxiety showed strong associations between RPA and social functioning. RPA was linked to better social functioning in males, but worse social functioning in mood and anxiety. This effect was especially pronounced for mood sufferers (43 total effects). Two effects in DRD4-7R carriers and 12 in non-right handed individuals indicated RPA association to self-oriented, rather than socially oriented, characteristics. For DRD4-7R carriers greater RPA was linked to increased self-determination and reduced novelty seeking (positive outcomes). For non-right handed individuals, greater RPA was linked to increased novelty seeking and reduced persistence and self-determination (negative outcomes). The locations of temperament effects were generally mixed, except for males (all P8-7), and non-right-hander (mainly at the TP8-7). Finally, of 31 findings involving the CPT, all but three were linked to social functioning, suggesting that this task taps brain-function characteristics important for social operations (Summary **Table 10**). For result details on the association between

TABLE 10 | Summary of RPA association with temperament by ADHD risk factor type.

Assessments	DRD4 7R	Males	Mood	Anxiety	Non right-handed
RPA association to temperament	Freq/Loc (2 eff) All Alpha P8-7 (1) P4-3 (1) Findings Less NS (1) More SD (1)	Freq/Loc (7 eff) Beta (6) P8-7 (7) Findings Less RD (1) Less CP (1) More SD (1) Better social (4)	Freq/Loc (43 eff) Alpha (29) TP8-7 (10) P8-7 (21) P4-3 (12) Findings More CP (2) More ST (2) Worse social (39)	Freq/Loc (6 eff) Alpha (4) TP8-7 (1) P8-7 (3) P4-3 (2) Findings Less RD (1) Less SD (1) More HA (1) Worse social (3)	Freq/Loc (12 eff) Alpha (9) TP8-7 (8) P8-7 (3) P4-3 (1) Findings More NS (2) Less PS (6) Less SD (4)

For "RPA association to temperament" more RPA (i.e., greater rightward parietal asymmetry) is associated with the reported outcomes; values in parentheses indicate the number of results (i.e., comprised of different frequency bands and/or parietal locations) showing a given effect. See **Table 2** for description of abbreviations; Freq/Loc, frequency/location; eff, effects.

TABLE 11 | Summary of RPA Association with Cognition by ADHD Risk Factor Type.

Assessments	DRD4 7R	Males	Mood	Anxiety	Non Right-handed
RPA association to cognition	Freq/Loc (15 eff) Alpha (11) TP8-7 (8) P8-7 (3) P4-3 (4) Better (8 eff) Read_rec (2) St-Word (1) Arith (2) SWM RTSD (3) Worse (7 eff) Omiss (3) Hit RTSE (1) Bias (3)	Freq/Loc (18 eff) Beta (16) TP8-7 (7) P8-7 (3) P4-3 (8) Better (18 eff) Read_rec (1) DSF-acc/max (3) SSF-acc/max (5) SSB-acc (1) SWM Acc (3) SWM RTSD (3) Hit RTSE (1) D-prime (1)	Freq/Loc (13 eff) All Beta All TP8 Better (13 eff) St-Word (1) SWM Acc (5) SWM RTSD (1) Omiss (2) D-prime (3) Bias (1)	Freq/Loc (15 eff) Beta (9) TP8-7 (6) P8-7 (8) P4-3 (1) Better (3 eff) SSF-acc (1) SWM RTSD (2) Worse (12 eff) Read_rec (2) Phono (1) Coding (1) St-Inter (4) DSF-acc/max (3) SSB-max (1)	Freq/Loc (29 eff) Beta (24) TP8-7 (12) P8-7 (15) P4-3 (1) Better (15 eff) DSB-acc (1) SWM Acc (14) Worse (14 eff) ST-Inter (1) DSB-max (1) SWM RT (12)

For "RPA association to cognition" more RPA (i.e., greater rightward parietal asymmetry) is associated with the reported outcomes; Values in parentheses indicate the number of results (i.e., comprised of different frequency bands and/or parietal locations) showing a given effect. See **Table 2** for description of measures abbreviations; Freq/Loc, frequency/location; eff, effects.

RPA and temperament characteristics, see supplement Table S1 in Supplementary Material.

Parietal Asymmetry Association to Cognition in Different ADHD RFs

RPA was associated with cognitive abilities in all RFs. Males and mood showed only positive effects (i.e., RPA = better performance). DRD4-7R, anxiety, and non-right handed RFs showed both positive and negative effects. A link between greater RPA and better SWM ability was evident for all groups (plus VWM in non-right handers and males). In DRD4-7R carriers, greater RPA was associated with better linguistic, but poorer attention ability. In males and mood sufferers, greater RPA was associated with better linguistic and attention ability. In anxiety sufferers, RPA was associated worse linguistic, WM, and EF ability. In non-right-handers, it was associated with worse WM and EF ability.

Associations between RPA and cognitive abilities were most evident during active visual processing (CPT and EO conditions). Males showed abundant effects at the P4-3 LI. Mood only showed effects at the TP8-7 LI. Other RFs showed mixed associations to parietal LIs. In DRD4-7R carriers, RPA associations to cognition were mostly in the alpha frequencies. Anxiety showed mixed

frequency effects. Males, mood, and non-right-handers showed the majority of effects in beta frequencies. Lastly, CPT measured sustained attention was almost exclusively tied to RPA at the TP8-7 LI, while CPT and SWM tasks that required sustained attention and task-directed visual processing were predominantly linked to beta frequencies (Summary **Table 11**). For result details on the association between RPA and cognition, see Supplement Table S2 in Supplementary Material.

Part 2 Discussion

The DEP model of ADHD highlights that multiple distributed brain systems get dynamically coordinated during task-directed thought and action. We posited that these systems comprise an emergent task-directed neural architecture (i.e., the TD-APS) that mobilizes RH resources in a specific task-specialized manner that facilitates fast (i.e., economical) identification of task-relevant content. Any impairment to this system, no matter the cause, is expected to reduce this capacity, and produce an associated increased expression of RH attention and non-task-entrained visuo-perceptual processing. Variable deficits to this system are expected to underlie ADHD heterogeneity, and manifest variant forms of this shared characteristic. The model also postulates

that variable primary deficits might be elucidated by examining the relationship between RH-biased processing and constituent system operations. We explored this possibility in our secondary analyses.

As predicted, different ADHD RFs showed variant expressed forms of RPA, along with different patterns of association between RPA and clinical and cognitive metrics. This supports the view that RPA is tied to different mechanisms in different ADHD RFs. The current study design cannot resolve what these are; however, through consideration of our total study findings, we can begin to speculate about possible differences in this regard. The following discussion of this topic and associated findings is highly speculative, but is provided to help stimulate further conceptual and experimental analysis of this topic.

The most obvious difference across ADHD RFs was that DRD4-7R and males exhibited RPA in multiple brain regions, frequencies, and conditions, whereas mood and anxiety showed very specific effects. One possible explanation is that genetically linked risk for ADHD (DRD4-7R carriers and males) is tied to more generalized and stable brain-function characteristics, while more experience-linked conditions (mood and anxiety) involve more state and/or circumstance specific abnormal brain functioning. Regardless, this pattern indicates that categorically different forms of brain function may underlie RPA in more genetically versus experience weighted conditions.

Within these domains, additional unique RPA characteristics were evident. The DRD4-7R group was predominantly associated with RPA in the alpha band (11/18 effects in alpha), while being male was predominantly associated with RPA in the beta band (10/15 effects in beta). According to recent conceptualizations of these frequencies, this might indicate increased RH internal-processing control in DRD4-7R carriers (121–123), but greater expression of RH visuo-perceptual functioning in males [for review, see Ref. (124–126)].

Behavioral laterality outcomes appear to support this speculation. DRD4-7R carriers showed an atypical pattern of being faster for word targets in the right versus left hemisphere (111, 127). They also showed faster reaction times for a specific dichotic listening trial type that is understood to require left-to-right callosal transfer of information due to exclusive RH competence for the stimuli (i.e., detecting the negative tone of voice in the right-ear projecting initially to the LH). Both results are consistent with greater RH processing control in DRD4-7R carriers (111). In contrast, being male was associated with faster responses for the trial type that directly taps RH ability (i.e., detecting the negative tone of voice in the left-ear projecting directly to the RH). In sum, both RPA and behavioral laterality findings seem consistent with greater RH processing control in DRD4-7R carriers, but greater expression of RH visuo-perceptual functions in males.

Mood and anxiety also differed in their expression of RPA and behavioral laterality outcomes. In mood, RPA was only evident during the EC condition (with no active sensory engagement). In anxiety, it was only evident during the EO and CPT conditions (with active sensory engagement). Consistent with this, mood showed no behavioral laterality effects during the dichotic listening task that requires active sensory processing, while anxiety was associated with faster responses for the trial type (noted above)

that requires left-to-right transfer of information (111). Hence, RPA and behavioral laterality outcomes suggest that $R > L$ parietal brain function may only be evident in mood sufferers when they are not sensory-engaged, and in anxiety sufferers when they are. This pattern, along with the above described variant expressions of RPA in DRD4-7R carriers and males, generally supports the view that unique mechanisms may underlie poor task-directed brain function and RPA across different ADHD RFs.

Additional Specific Hypotheses

In order to generate more specific hypothesis about the nature of abnormal brain function across ADHD RFs, we will now consider individual RFs clinical characteristics, and patterns of RPA association to symptoms, temperament, and cognitive abilities. This analysis is again speculative, but is presented to help generate testable hypotheses for future analysis.

The DRD4-7R allele

Carrying the DRD4-7R allele was generally distinguished by having more positive outcomes, and RPA effects being mainly in the alpha band. It was associated with having fewer inattentive symptoms and an advantage for spatial working memory. RPA (mainly in alpha) was associated with fewer inattentive symptoms, positive temperament characteristics, and better ability for tasks amenable to visual strategies. Negative RPA associations were exclusive to the CPT, and indicated a link to impulsive and variable performance.

The CPT requires verbal encoding and maintaining a task-directed set for a prolonged period (i.e., 14 min). This perhaps makes the CPT less amenable to alternative RH strategies. Moreover, negative RPA-CPT performance associations only occurred for RPA in the TPJ (i.e., TP8-TP7). This may indicate a link between cognitive flexibility and poor CPT performance in this ADHD RF. Given this pattern, and indications of strong positive selection for the DRD4-7R allele (128), we speculate that this RF might reflect an evolved specialization for more flexible visual cognition that supports exploratory behavior, but is deleterious for task-directed actions.

Being male

Being male was generally distinguished by poorer verbal ability, and by RPA effects occurring mainly in the beta band. Males exhibited worse letter discrimination and coding ability. RPA (mainly in beta) was associated with greater hyperactivity, better social functioning, and better cognitive ability (e.g., spatial, verbal, and less response variability). As with the DRD4-7R carriers, we speculate that being male is linked to an evolved emphasis on RH-biased brain function that is less conducive to task-directed thought and behavior, albeit in a different manner.

EEG beta has been linked to early-stage encoding of attentionally selected information [for review, see Ref. (124–126)] and shows lateralized posterior expression with verbal and visual-spatial processing (129, 130). Hence, the emphasis of EEG beta and poorer verbal ability in males (74, 75) seems consistent with greater weighting of RH-biased sensory encoding. In this vein, greater RH visual-cortical volumes have been identified in males

(131), and abundant gender differences in visual processing are identified [for review, see Ref. (132, 133)].

Women exhibit greater specialization for top-down object-oriented processing (i.e., categorization/naming), while men emphasize bottom-up analysis of motion, fine detail, and high resolution spatial and temporal content [for review, see Ref. (132, 134)]. Moreover, a recent study has demonstrated that while the female style of sensory processing is well adapted to EF-demanding dual-task paradigms, the male style is not (132). Given this literature and our current findings, we speculate that risk for ADHD in males is linked to greater expression of RH bottom-up visual processing, which is generally less conducive to the induction and/or maintenance of a task-directed neural architecture. Perhaps consistent with this male RPA effects on cognition were wholly positive, including for CPT metrics, suggesting that compensatory attentional effort might be an important driver of RPA in males.

Having mood disorder

Having mood disorder was distinguished by greater inattention, poorer inhibition, RPA being exclusively expressed during the EC condition, and no RPA associations to symptoms, but abundant RPA associations to poor social functioning. For example, mood sufferers exhibited greater CPT commission errors (i.e., poor response inhibition). Poor inhibition is consistent with RH frontal pathology in mood disorders (57, 135). Moreover, mood was the only RF that did not exhibit RPA association to symptoms; however, they exhibited far more RPA associations to bad social outcomes than any other group. Finally, as in males, RPA associations to cognition were exclusively positive.

Taken together, we speculate that RH bias in mood sufferers, identified here and elsewhere [for review, see Ref. (135)], may reflect internal affective processing (i.e., rumination) that imparts reduced awareness of external cues needed to guide attention in a socially adaptive manner (i.e., poor entrainment of attention functions to external social contexts). Perhaps consistent with this, mood-linked RPA was only evident in SPL and IPL regions, which are often linked to applied attentional effects.

Furthermore, greater $R > L$ TPJ activation was tied to better cognitive outcomes for the mood RF. It is interesting to consider that a more flexible cognitive set might help mood sufferers be more responsive to external social cues, and thereby make socially appropriate transitions into task-directed cognitive sets. If true, mood derived risk for ADHD should be moderated by factors that promote more flexible and/or externally oriented cognitive and attentional sets (e.g., fluid intelligence, creative ability, self-transcendence). Consistent with this, in our ADHD sample, comorbid mood was associated with low self-transcendence (i.e., feeling less connected to the external environment) (136).

Having anxiety disorder

Having anxiety disorder was distinguished by being the most impaired RF, by RPA occurring only during sensory engagement, and by having the most numerous RPA associations to symptoms and poor cognition. The DRD4-7R group was associated with reduced general knowledge (information task), being male was associated with poorer verbal ability, and mood was associated with poor inhibition, having anxiety disorder was associated with

each. It was also tied to greater hyperactivity, harm avoidance, and social problems. RPA showed the most numerous associations with symptoms, marginal association to poor social functioning, and mixed effects on cognition, but with the most numerous negative effects. Different from DRD4-7R carriers, anxiety sufferers showed a standard pattern of RPA association to positive outcomes for spatial tasks, and poor outcomes for standard verbal tasks and the CPT. Hence, rather than increased RH processing control, we speculate that anxiety imparts a more fight-or-flight like increase of bottom-up visuo-perceptual processing. If true, anxiety may produce a similar, albeit more severe, version of the ADHD liability that we suggest occurs in males (i.e., greater expression of bottom-up visual processing).

In considering the sum total of our findings, along with relevant literature, we speculate that RPA is tied to different causal mechanisms in different ADHD RFs. We suspect that DRD4-7R carriers may adhere to visual cognition strategies that bear liability for tasks that are not amenable to visual cognition strategies. Males and anxiety sufferers may bear an increased tendency toward bottom-up visual processing (automatic sensory responsivity) that makes the induction and maintenance of a task-directed neural architecture more difficult (especially for anxiety sufferers). We suspect that in males this is typically overcome, but nevertheless imparts risk for ADHD, while in anxiety sufferers, it more consistently underlies attention related pathology. Finally, we speculated that internal affective processing in mood sufferers might diminish entrainment to social cues needed to guide attention.

General Summary and Conclusion

The DEP model conceptualizes ADHD to be a set of clinical characteristics that manifest with any form of poor task-directed brain functioning (i.e., convergent effects). It asserts that $R > L$ posterior brain function is a similarly manifest physiologic outcome. We tested this by examining whether $R > L$ parietal EEG asymmetry was generally associated with common ADHD RFs, and found that it was. This suggests that ADHD liability is generally associated with increased weighting of RH visuo-perceptual versus LH verbal parietal brain functions, and highlights two possible mechanisms: (1) a TPJ state-setting effect and (2) an SPL/IPL applied attention effect. Moreover, we hypothesize that abnormal RH contribution in ADHD during task operations can be conceived to include: (1) reduced top-down task-directed control over visual processing (low RH), (2) a more diffuse/flexible attentional and cognitive set, associated with greater expression of non-task-specialized forms of visual processing (high RH), and (3) compensatory attentional effort (high RH).

In secondary analyses, we explored whether RPA (a convergent feature of ADHD liability) might be tied to different causal mechanisms in different ADHD RFs. Different patterns of RPA and RPA association to symptoms, temperament, and cognition suggest that it is. We suspect that ADHD RFs disrupt the induction and/or maintenance of a task-optimized neural architecture (i.e., a TD-APS), albeit in different manners. We also expect that multiple additional acute (e.g., sleep deprivation, highly engaging sensory experience, emergencies, etc.), developmental (e.g., childhood trauma/abuse), genetic (e.g., that promote a novelty/risk-taking temperament), and/or mechanistic (e.g., brain

injury, learning disabilities) factors can produce this outcome. Under this view, ADHD symptoms are considered to reflect the convergent or shared outcome of poor task-directed brain functioning, no matter the cause, with the syndrome of ADHD reflecting such circumstances that get durably expressed and that are life disruptive.

Viewed in this way, it is clear that the challenge of understanding and treating ADHD is, at its core, synonymous with the challenge of understanding complex task-directed thought and action, and the multitude of ways that ability might be compromised. It also highlights that identification of primary deficits may require a greater emphasis on longitudinal analysis of ADHD individuals and/or clinically meaningful subgroups. However, as demonstrated by current drug treatments, alleviation of ADHD symptoms may not require a complete knowledge of deficit causality.

Given the diffuse anatomy of catecholamine systems, stimulants likely impart broad, rather than specific effects (137). One possibility is that they help to generally nudge brain function toward computational states that are conducive to task-directed thought and action, and thus provide a degree of clinical benefit regardless of diverse deficit sources. Moreover, the two primary systems targeted by ADHD drug therapies, dopamine and norepinephrine, appear to exhibit a degree of left and right hemisphere specialization (77, 138), which may also align with identified left–right dichotomies related to internal-verbal versus external-perceptual brain functions (129). It is interesting to consider that stimulants might help to achieve a task-optimized leftward balance along these continuums (i.e., greater brain-state orientation toward internal-verbal versus external-perceptual processing). If true, additional non-drug based therapies aimed at bolstering verbal and internally weighted brain function may prove helpful for ADHD individuals. To this end, we recently used transcranial magnetic stimulation (TMS) to increase relative LH parietal contribution (near TPJ) in ADHD adults, and obtained highly

encouraging preliminary results. We think that cognitive training strategies previously shown to increase bias toward LH verbal-categorical processing [for review, see Ref. (139)] should also be explored.

In sum, the current study and DEP model of ADHD have attempted to highlight, and begin to address, the challenges associated with trying to map sharply defined diagnostic categories onto underlying pathophysiology, as abnormal brain-function characteristics are often shared by multiple disorders. To this end, aberrant rightward asymmetry has been reported across multiple conditions that are linked to attention difficulties, and that are often comorbid with ADHD (e.g., anxiety, depression, sleep deprivation, novelty seeking, reading disability, etc.) (8, 131, 132, 140–147). Our core assertion is that this phenotype is generally linked to brain-function dynamics that are not optimized for complex multi-step and/or contingency-based task-directed sensory processing and cognition. Accordingly, and as noted, we expect that the effects observed in the current study can occur independent of an ADHD diagnosis; however, and importantly, we do not expect them to be independent of attentional liability for complex task-directed brain-function. Whether or not an ADHD diagnosis gets applied is a matter of clinical judgment.

Acknowledgments

This work was funded in part by National Institute of Mental Health Grant MH058277 (SS), National Institute of Child Health and Human Development Grant HD40275 (SL), National Institute of Neurological Disease and Stroke NS054124 (SL), and National Institute of Mental Health Grant MH082104 (TSH).

Supplementary Material

The Supplementary Material for this article can be found online at <http://journal.frontiersin.org/article/10.3389/fpsy.2015.00063>

References

- Hale TS. A distributed effects perspective of dimensionally defined psychiatric disorders: and convergent versus core deficit effects in ADHD. *Front Psychiatry* (2014) 5:62. doi:10.3389/fpsy.2014.00062
- Heilman KM, Bowers D, Valenstein E, Watson RT. The right hemisphere: neuropsychological functions. *J Neurosurg* (1986) 64(5):693–704. doi:10.3171/jns.1986.64.5.0693
- Ruff CC, Blankenburg F, Boertomto O, Bestmann S, Weiskopf N, Driver J. Hemispheric differences in frontal and parietal influences on human occipital cortex: direct confirmation with concurrent TMS-fMRI. *J Cogn Neurosci* (2009) 21(6):1146–61. doi:10.1162/jocn.2009.21097
- Zanto TP, Rubens MT, Thangavel A, Gazzaley A. Causal role of the prefrontal cortex in top-down modulation of visual processing and working memory. *Nat Neurosci* (2011) 14(5):656–61. doi:10.1038/nn.2773
- Summerfield C, Egner T. Expectation (and attention) in visual cognition. *Trends Cogn Sci* (2009) 13(9):403–9. doi:10.1016/j.tics.2009.06.003
- Pardo JV, Raichle ME. Localization of a human system of sustained attention by positron emission tomography. *Nature* (1991) 349:61–4. doi:10.1038/349061a0
- Corbetta M, Miezin F, Shulman G, Petersen S. A PET study of visuospatial attention. *J Neurosci* (1993) 13(3):1202–26.
- Bubic A, von Cramon DY, Schubotz RI. Exploring the detection of associatively novel events using fMRI. *Hum Brain Mapp* (2011) 32(3):370–81. doi:10.1002/hbm.21027
- Downar J, Crawley AP, Mikulis DJ, Davis KD. The effect of task relevance on the cortical response to changes in visual and auditory stimuli: an event-related fMRI study. *Neuroimage* (2001) 14(6):1256–67. doi:10.1006/nimg.2001.0946
- Roser ME, Fiser J, Aslin RN, Gazzaniga MS. Right hemisphere dominance in visual statistical learning. *J Cogn Neurosci* (2011) 23(5):1088–99. doi:10.1162/jocn.2010.21508
- Zeithamova D, Maddox WT. The role of visuospatial and verbal working memory in perceptual category learning. *Mem Cognit* (2007) 35(6):1380–98. doi:10.3758/BF03193609
- Di Dio C, Macaluso E, Rizzolatti G. The golden beauty: brain response to classical and renaissance sculptures. *PLoS One* (2007) 2(11):e1201. doi:10.1371/journal.pone.0001201
- Hammer R, Brechmann A, Ohl F, Weinshall D, Hochstein S. Differential category learning processes: the neural basis of comparison-based learning and induction. *Neuroimage* (2010) 52(2):699–709. doi:10.1016/j.neuroimage.2010.03.080
- Hochstein S, Ahissar M. View from the top: hierarchies and reverse hierarchies in the visual system. *Neuron* (2002) 36(5):791–804. doi:10.1016/S0896-6273(02)01091-7
- Corbetta M, Shulman GL. Control of goal-directed and stimulus-driven attention in the brain. *Nat Rev Neurosci* (2002) 3(3):201–15. doi:10.1038/nrn755
- Bar M. A cortical mechanism for triggering top-down facilitation in visual object recognition. *J Cogn Neurosci* (2003) 15(4):600–9. doi:10.1162/089892903321662976

17. Hale TS, Loo SK, Zaidel E, Hanada G, Macion J, Smalley SL. Rethinking a right hemisphere deficit in ADHD. *J Atten Disord* (2009) **13**(1):3–17. doi:10.1177/1087054708323005
18. Hale TS, McCracken JT, McGough JJ, Smalley SL, Phillips JM, Zaidel E. Impaired linguistic processing and atypical brain laterality in adults with ADHD. *Clin Neurosci Res* (2005) **5**:255–63. doi:10.1016/j.neuropsychologia.2010.08.002
19. Hale TS, Zaidel E, McGough JJ, Phillips JM, McCracken JT. Atypical brain laterality in adults with ADHD during dichotic listening for emotional intonation and words. *Neuropsychologia* (2006) **44**(6):896–904. doi:10.1016/j.neuropsychologia.2005.08.014
20. Hale TS, Bookheimer S, McGough JJ, Phillips JM, McCracken JT. Atypical brain activation during simple & complex levels of processing in adult ADHD: an fMRI study. *J Atten Disord* (2007) **11**(2):125–40. doi:10.1177/1087054706294101
21. Hale TS, Kane AM, Kaminsky O, Tung KL, Wiley JF, McGough JJ, et al. Visual network asymmetry and default mode network function in ADHD: an fMRI study. *Front Psychiatry* (2014) **5**:81. doi:10.3389/fpsy.2014.00081
22. Hale TS, Kane AM, Tung KL, Kaminsky O, McGough JJ, Hanada G, et al. Abnormal parietal brain function in ADHD: replication and extension of previous EEG beta asymmetry findings. *Front Psychiatry* (2014) **5**:87. doi:10.3389/fpsy.2014.00087
23. Hale TS, Smalley SL, Dang J, Hanada G, Macion J, McCracken JT, et al. ADHD familial loading and abnormal EEG alpha asymmetry in children with ADHD. *J Psychiatr Res* (2010) **44**(9):605–15. doi:10.1016/j.jpsychires.2009.11.012
24. Hale TS, Smalley SL, Hanada G, Macion J, McCracken JT, McGough JJ, et al. Atypical alpha asymmetry in adults with ADHD. *Neuropsychologia* (2009) **47**(10):2082–8. doi:10.1016/j.neuropsychologia.2009.03.021
25. Hale TS, Smalley SL, Walshaw PD, Hanada G, Macion J, McCracken JT, et al. Atypical EEG beta asymmetry in adults with ADHD. *Neuropsychologia* (2010) **48**(12):3532–9. doi:10.1016/j.neuropsychologia.2010.08.002
26. Clarke AR, Barry RJ, Heaven PC, McCarthy R, Selikowitz M, Byrne MK. EEG in adults with attention-deficit/hyperactivity disorder. *Int J Psychophysiol* (2008) **70**(3):176–83. doi:10.1016/j.ijpsycho.2008.07.001
27. Clarke AR, Barry RJ, McCarthy R, Selikowitz M. EEG analysis of children with attention-deficit/hyperactivity disorder and comorbid reading disabilities. *J Learn Disabil* (2002) **35**(3):276–85. doi:10.1177/002221940203500309
28. Cortese S, Kelly C, Chabernaud C, Proal E, Di Martino A, Milham MP, et al. Toward systems neuroscience of ADHD: a meta-analysis of 55 fMRI studies. *Am J Psychiatry* (2012) **169**(10):1038–55. doi:10.1176/appi.ajp.2012.11101521
29. Johnstone SJ, Barry RJ, Clarke AR. Ten years on: a follow-up review of ERP research in attention-deficit/hyperactivity disorder. *Clin Neurophysiol* (2013) **124**(4):644–57. doi:10.1016/j.clinph.2012.09.006
30. Willcutt EG, Doyle AE, Nigg JT, Faraone SV, Pennington BF. Validity of the executive function theory of attention-deficit/hyperactivity disorder: a meta-analytic review. *Biol Psychiatry* (2005) **57**(11):1336. doi:10.1016/j.biopsych.2005.02.006
31. Weiler MD, Bernstein JH, Bellinger DC, Waber DP. Processing speed in children with attention deficit hyperactivity disorder, inattentive type. *Child Neuropsychol* (2000) **6**(3):218–34. doi:10.1076/chin.6.3.218.3156
32. Tannock R, Martinussen R, Frijters J. Naming speed performance and stimulant effects indicate effortful, semantic processing deficits in attention-deficit/hyperactivity disorder. *J Abnorm Child Psychol* (2000) **28**(3):237–52. doi:10.1023/A:1005192220001
33. Stevens J, Quittner AL, Zuckerman JB, Moore S. Behavioral inhibition, self-regulation of motivation, and working memory in children with attention deficit hyperactivity disorder. *Dev Neuropsychol* (2002) **21**(2):117–39. doi:10.1207/S15326942DN2102_1
34. Semrud-Clikeman M, Guy K, Griffin JD, Hynd GW. Rapid naming deficits in children and adolescents with reading disabilities and attention deficit hyperactivity disorder. *Brain Lang* (2000) **75**(1):70–83. doi:10.1006/brln.2000.2337
35. Rucklidge JJ, Tannock R. Neuropsychological profiles of adolescents with ADHD: effects of reading difficulties and gender. *J Child Psychol Psychiatry* (2002) **43**(8):988–1003. doi:10.1111/1469-7610.00227
36. Nigg JT, Butler KM, Huang-Pollock CL, Henderson JM. Inhibitory processes in adults with persistent childhood onset ADHD. *J Consult Clin Psychol* (2002) **70**(1):153–7. doi:10.1037/0022-006X.70.1.153
37. Brock SE, Christo C. Digit naming speed performance among children with attention-deficit/hyperactivity disorder. *Calif Sch Psychol* (2003) **8**:115–25. doi:10.1007/BF03340900
38. Malone MA, Kershner JR, Siegel L. The effects of methylphenidate on levels of processing and laterality in children with attention deficit disorder. *J Abnorm Child Psychol* (1988) **16**(4):379–95. doi:10.1007/BF00914170
39. Cohen MS, Kosslyn SM, Breiter HC, DiGirolamo GJ, Thompson WL, Bookheimer SY, et al. Changes in cortical activity during mental rotation: a mapping study using functional magnetic resonance imaging. *Brain* (1996) **119**:89–100. doi:10.1093/brain/119.1.89
40. Ernst M, Zametkin AJ, Matochik JA, Jons PH, Cohen RM. DOPA decarboxylase activity in attention deficit hyperactivity disorder adults. A [fluorine-18]fluorodopa positron emission tomographic study. *J Neurosci* (1998) **18**(15):5901–7.
41. Seig KG, Gaffney GR, Preston DF, Jellings JA. SPECT brain imaging abnormalities in attention deficit hyperactivity disorder. *Clin Nucl Med* (1995) **20**:55–60. doi:10.1097/00003072-199501000-00014
42. Zametkin AJ, Nordahl TE, Gross M, King AC, Semple WE, Rumsy J, et al. Cerebral glucose metabolism in adults with hyperactivity of childhood onset. *N Engl J Med* (1990) **323**:1361–6. doi:10.1056/NEJM199011153232001
43. Zametkin AJ, Liebenauer LL, Fitzgerald GA, King AC, Minkunas DV, Herscovitch P, et al. Brain metabolism in teenagers with attention-deficit hyperactivity disorder. *Arch Gen Psychiatry* (1993) **50**(5):333–40. doi:10.1001/archpsyc.1993.01820170011002
44. Baving L, Laucht M, Schmidt MH. Atypical frontal brain activation in ADHD: preschool and elementary school boys and girls. *J Am Acad Child Adolesc Psychiatry* (1999) **38**(11):1363–71. doi:10.1097/00004583-199911000-00010
45. Chabot RJ, Serfontein G. Quantitative electroencephalographic profiles of children with attention deficit disorder. *Biol Psychiatry* (1996) **40**:951–63. doi:10.1016/0006-3223(95)00576-5
46. Swartwood JN, Swartwood MO, Lubar JF, Timmermann DL. EEG differences in ADHD-combined type during baseline and cognitive tasks. *Pediatr Neurol* (2003) **28**(3):199–204. doi:10.1016/S0887-8994(02)00514-3
47. Silk TJ, Vance A, Rinehart N, Bradshaw JL, Cunnington R. White-matter abnormalities in attention deficit hyperactivity disorder: a diffusion tensor imaging study. *Hum Brain Mapp* (2009) **30**(9):2757–65. doi:10.1002/hbm.20703
48. Li Q, Sun J, Guo L, Zang Y, Feng Z, Huang X, et al. Increased fractional anisotropy in white matter of the right frontal region in children with attention-deficit/hyperactivity disorder: a diffusion tensor imaging study. *Neuro Endocrinol Lett* (2010) **31**(6):747–53.
49. Li X, Jiang J, Zhu W, Yu C, Sui M, Wang Y, et al. Asymmetry of prefrontal cortical convolution complexity in males with attention-deficit/hyperactivity disorder using fractal information dimension. *Brain Dev* (2007) **29**(10):649–55. doi:10.1016/j.braindev.2007.04.008
50. Wang J, Jiang T, Cao Q, Wang Y. Characterizing anatomic differences in boys with attention-deficit/hyperactivity disorder with the use of deformation-based morphometry. *AJNR Am J Neuroradiol* (2007) **28**(3):543–7.
51. Seidman LJ, Valera EM, Makris N. Structural brain imaging of attention-deficit/hyperactivity disorder. *Biol Psychiatry* (2005) **57**(11):1263–72. doi:10.1016/j.biopsych.2004.11.019
52. Rolfé MH, Kirk IJ, Waldie KE. Interhemispheric callosal transfer in adults with attention-deficit/hyperactivity disorder: an event-related potential study. *Neuroreport* (2007) **18**(3):255–9. doi:10.1097/WNR.0b013e328011e6f9
53. Barry RJ, Clarke AR, McCarthy R, Selikowitz M, Johnstone SJ. EEG coherence adjusted for inter-electrode distance in children with attention-deficit/hyperactivity disorder. *Int J Psychophysiol* (2005) **58**(1):12–20. doi:10.1016/j.ijpsycho.2005.03.005
54. Clarke AR, Barry RJ, McCarthy R, Selikowitz M, Johnstone SJ, Hsu CI, et al. Coherence in children with attention-deficit/hyperactivity disorder and excess beta activity in their EEG. *Clin Neurophysiol* (2007) **118**(7):1472–9. doi:10.1016/j.clinph.2007.04.006
55. Banich MT, Burgess GC, Depue BE, Ruzic L, Bidwell LC, Hitt-Laustsen S, et al. The neural basis of sustained and transient attentional control in young adults with ADHD. *Neuropsychologia* (2009) **47**(14):3095–104. doi:10.1016/j.neuropsychologia.2009.07.005
56. Cubillo A, Halari R, Smith A, Taylor E, Rubia K. A review of fronto-striatal and fronto-cortical brain abnormalities in children and adults with attention deficit hyperactivity disorder (ADHD) and new evidence for dysfunction in adults

- with ADHD during motivation and attention. *Cortex* (2012) **48**(2):194–215. doi:10.1016/j.cortex.2011.04.007
57. Rubia K, Cubillo A, Smith AB, Woolley J, Heyman I, Brammer MJ. Disorder-specific dysfunction in right inferior prefrontal cortex during two inhibition tasks in boys with attention-deficit hyperactivity disorder compared to boys with obsessive-compulsive disorder. *Hum Brain Mapp* (2010) **31**(2):287–99. doi:10.1002/hbm.20864
 58. Tamm L, Menon V, Reiss AL. Parietal attentional system aberrations during target detection in adolescents with attention deficit hyperactivity disorder: event-related fMRI evidence. *Am J Psychiatry* (2006) **163**(6):1033–43. doi:10.1176/appi.ajp.163.6.1033
 59. Casey BJ, Castellanos FX, Geidd JN, Marsh WL, Hamburger SD, Schubert AB, et al. Implications of right frontostriatal circuitry in response inhibition and attention-deficit/hyperactivity disorder. *J Am Acad Child Adolesc Psychiatry* (1997) **36**(3):374–83. doi:10.1097/00004583-199703000-00016
 60. Casey BJ, Epstein JN, Buhle J, Liston C, Davidson MC, Tonev ST, et al. Frontostriatal connectivity and its role in cognitive control in parent-child dyads with ADHD. *Am J Psychiatry* (2007) **164**(11):1729–36. doi:10.1176/appi.ajp.2007.06101754
 61. Hill DE, Yeo RA, Campbell RA, Hart B, Vigil J, Brooks W. Magnetic resonance imaging correlates of attention-deficit/hyperactivity disorder in children. *Neuropsychology* (2003) **17**(3):496–506. doi:10.1037/0894-4105.17.3.496
 62. Yeo RA, Hill DE, Campbell RA, Vigil J, Petropoulos H, Hart B, et al. Proton magnetic resonance spectroscopy investigation of the right frontal lobe in children with attention-deficit/hyperactivity disorder. *J Am Acad Child Adolesc Psychiatry* (2003) **42**(3):303–10. doi:10.1097/00004583-200303000-00010
 63. Mataró M, García-Sánchez C, Junqué C, Estévez-González A, Pujol J. Magnetic resonance imaging measurements of the caudate nucleus in adolescents with attention-deficit hyperactivity disorder and its relationship with neuropsychological and behavioral measures. *Arch Neurol* (1997) **54**:963–8. doi:10.1001/archneur.1997.00550200027006
 64. Ernst M, Kimes AS, London ED, Matochik JA, Eldreth D, Tata S, et al. Neural substrates of decision making in adults with attention deficit hyperactivity disorder. *Am J Psychiatry* (2003) **160**(6):1061–70. doi:10.1176/appi.ajp.160.6.1061
 65. Vaidya CJ, Bunge SA, Dudukovic NM, Zalecki CA, Elliott GR, Gabrieli JD. Altered neural substrates of cognitive control in childhood ADHD: evidence from functional magnetic resonance imaging. *Am J Psychiatry* (2005) **162**(9):1605–13. doi:10.1176/appi.ajp.162.9.1605
 66. Burgess GC, Depue BE, Ruzic L, Willcutt EG, Du YP, Banich MT. Attentional control activation relates to working memory in attention-deficit/hyperactivity disorder. *Biol Psychiatry* (2010) **67**(7):632–40. doi:10.1016/j.biopsych.2009.10.036
 67. Keune PM, Schönberg M, Wyckoff S, Mayer K, Riemann S, Hautzinger M, et al. Frontal alpha-asymmetry in adults with attention deficit hyperactivity disorder: replication and specification. *Biol Psychol* (2011) **87**(2):306–10. doi:10.1016/j.biopsycho.2011.02.023
 68. Fassbender C, Schweitzer JB. Is there evidence for neural compensation in attention deficit hyperactivity disorder? A review of the functional neuroimaging literature. *Clin Psychol Rev* (2006) **26**(4):445–65. doi:10.1016/j.cpr.2006.01.003
 69. Reid HM, Norvilitis JM. Evidence for anomalous laterization across domain in ADHD children as well as adults identified with the wender Utah rating scale. *J Psychiatr Res* (2000) **34**:311–6. doi:10.1016/S0022-3956(00)00027-3
 70. Niederhofer H. Hand preference in attention deficit hyperactivity disorder. *Percept Mot Skills* (2005) **101**(3):808–10. doi:10.2466/pms.101.3.808-810
 71. Toga AW, Thompson PM. Mapping brain asymmetry. *Nat Rev Neurosci* (2003) **4**(1):37–48. doi:10.1038/nrn1009
 72. Geschwind DH, Miller BL, DeCarli C, Carmelli D. Heritability of lobar brain volumes in twins supports genetic models of cerebral laterality and handedness. *Proc Natl Acad Sci U S A* (2002) **99**(5):3176–81. doi:10.1073/pnas.052494999
 73. Berry CA, Shaywitz SE, Shaywitz BA. Girls with attention deficit disorder: a silent minority? A report on behavioral and cognitive characteristics. *Pediatrics* (1985) **76**(5):801–9.
 74. Jones CM, Braithwaite VA, Healy SD. The evolution of sex differences in spatial ability. *Behav Neurosci* (2003) **117**(3):403–11. doi:10.1037/0735-7044.117.3.403
 75. Joseph R. The evolution of sex differences in language, sexuality, and visual-spatial skills. *Arch Sex Behav* (2000) **29**(1):35–66. doi:10.1023/A:1001834404611
 76. Pliszka SR. The neuropsychopharmacology of attention-deficit/hyperactivity disorder. *Biol Psychiatry* (2005) **57**(11):1385–90. doi:10.1016/j.biopsych.2004.08.026
 77. Tucker DM, Williamson PA. Asymmetric neural control systems in human self-regulation. *Psychol Rev* (1984) **91**(2):185–215. doi:10.1037/0033-295X.91.2.185
 78. Bressler SL, Tognoli E. Operational principles of neurocognitive networks. *Int J Psychophysiol* (2006) **60**(2):139–48. doi:10.1016/j.ijpsycho.2005.12.008
 79. Baddeley A, Chincotta D, Adlam A. Working memory and the control of action: evidence from task switching. *J Exp Psychol Gen* (2001) **130**(4):641–57. doi:10.1037/0096-3445.130.4.641
 80. Desimone R, Duncan J. Neural mechanisms of selective visual attention. *Annu Rev Neurosci* (1995) **18**:193–222. doi:10.1146/annurev.ne.18.030195.001205
 81. Olivers CN, Peters J, Houtkamp R, Roelofsma PR. Different states in visual working memory: when it guides attention and when it does not. *Trends Cogn Sci* (2011) **15**(7):327–34. doi:10.1016/j.tics.2011.05.004
 82. de Fockert JW. Beyond perceptual load and dilution: a review of the role of working memory in selective attention. *Front Psychol* (2013) **4**:287. doi:10.3389/fpsyg.2013.00287
 83. Ghajar J, Ivry RB. The predictive brain state: asynchrony in disorders of attention? *Neuroscientist* (2009) **15**(3):232–42. doi:10.1177/1073858408326429
 84. Kaller CP, Rahm B, Spreer J, Weiller C, Unterrainer JM. Dissociable contributions of left and right dorsolateral prefrontal cortex in planning. *Cereb Cortex* (2011) **21**(2):307–17. doi:10.1093/cercor/bhq096
 85. Hickok G, Poeppel D. The cortical organization of speech processing. *Nat Rev Neurosci* (2007) **8**(5):393–402. doi:10.1038/nrn2113
 86. Barkley RA. Behavioral inhibition, sustained attention, and executive functions: construction a unifying theory of ADHD. *Psychol Bull* (1997) **121**:65–94. doi:10.1037/0033-2909.121.1.65
 87. Alderson RM, Rapport MD, Hudec KL, Sarver DE, Kofler MJ. Competing core processes in attention-deficit/hyperactivity disorder (ADHD): do working memory deficiencies underlie behavioral inhibition deficits? *J Abnorm Child Psychol* (2010) **38**(4):497–507. doi:10.1007/s10802-010-9387-0
 88. Hoekzema E, Carmona S, Ramos-Quiroga JA, Richarte Fernandez V, Bosch R, Soliva JC, et al. An independent components and functional connectivity analysis of resting state fMRI data points to neural network dysregulation in adult ADHD. *Hum Brain Mapp* (2014) **35**(4):1261–72. doi:10.1002/hbm.22250
 89. Jacobson LA, Ryan M, Martin RB, Ewen J, Mostofsky SH, Denckla MB, et al. Working memory influences processing speed and reading fluency in ADHD. *Child Neuropsychol* (2011) **17**(3):209–24. doi:10.1080/09297049.2010.532204
 90. Rogers M, Hwang H, Toplak M, Weiss M, Tannock R. Inattention, working memory, and academic achievement in adolescents referred for attention deficit/hyperactivity disorder (ADHD). *Child Neuropsychol* (2011) **17**(5):444–58. doi:10.1080/09297049.2010.544648
 91. Castellanos FX, Sonuga-Barke EJ, Milham MP, Tannock R. Characterizing cognition in ADHD: beyond executive dysfunction. *Trends Cogn Sci* (2006) **10**(3):117–23. doi:10.1016/j.tics.2006.01.011
 92. Faraone SV, Mick E. Molecular genetics of attention deficit hyperactivity disorder. *Psychiatr Clin North Am* (2010) **33**(1):159–80. doi:10.1016/j.psc.2009.12.004
 93. Yoshimasu K, Barbaresi WJ, Colligan RC, Voigt RG, Killian JM, Weaver AL, et al. Childhood ADHD is strongly associated with a broad range of psychiatric disorders during adolescence: a population-based birth cohort study. *J Child Psychol Psychiatry* (2012) **53**(10):1036–43. doi:10.1111/j.1469-7610.2012.02567.x
 94. Smalley SL, McGough JJ, Del'Homme M, NewDelman J, Gordon E, Kim T, et al. Familial clustering of symptoms and disruptive behaviors in multiplex families with attention-deficit/hyperactivity disorder. *J Am Acad Child Adolesc Psychiatry* (2000) **39**(9):1135–43. doi:10.1097/00004583-200009000-00013
 95. Loo SK, Hale ST, Hanada G, Macion J, Shrestha A, McGough JJ, et al. Familial clustering and DRD4 effects on electroencephalogram measures in multiplex families with attention deficit/hyperactivity disorder. *J Am Acad Child Adolesc Psychiatry* (2010) **49**(4):368–77. doi:10.1016/j.jaac.2010.01.002
 96. Kaufman J, Birmaher B, Brent D, Rao U, Flynn C, Moreci P, et al. Schedule for affective disorders and schizophrenia for school-age children-present and lifetime version (K-SADS-PL): initial reliability and validity

- data. *J Am Acad Child Adolesc Psychiatry* (1997) **36**(7):980–8. doi:10.1097/00004583-199707000-00021
97. Berument SK, Rutter M, Lord C, Pickles A, Bailey A. Autism screening questionnaire: diagnostic validity. *Br J Psychiatry* (1999) **175**:444–51. doi:10.1192/bjp.175.5.444
 98. Leckman JF, Sholomskas D, Thompson WD, Belanger A, Weissman MM. Best estimate of lifetime psychiatric diagnosis: a methodological study. *Arch Gen Psychiatry* (1982) **39**(8):879–83. doi:10.1001/archpsyc.1982.04290080001001
 99. Kustanovich V, Ishii J, Crawford L, Yang M, McGough JJ, McCracken JT, et al. Transmission disequilibrium testing of dopamine-related candidate gene polymorphisms in ADHD: confirmation of association of ADHD with DRD4 and DRD5. *Mol Psychiatry* (2004) **9**(7):711–7. doi:10.1038/sj.mp.4001466
 100. Oldfield RC. Assessment and analysis of handedness: the Edinburgh inventory. *Neuropsychologia* (1971) **19**:97–113. doi:10.1016/0028-3932(71)90067-4
 101. Conners KC, Staff M. *Conners' Continuous Performance Test (CPT II)*. North Tonawanda, NY: Multi-Health Systems Inc (2002). 168 p.
 102. Conners CK, editor. *The Continuous Performance Test (CPT): Use as a Diagnostic Tool and Measure of Treatment Outcome*. Los Angeles, CA: American Psychological Association (1994).
 103. Welch P. The use of fast Fourier transform for the estimation of power spectra: a method based on time averaging over short, modified periodograms. *IEEE Trans Audio Electroacoust* (1967) **15**(2):70–3. doi:10.1109/TAU.1967.1161901
 104. Achenbach TM, Dumenci L. Advances in empirically based assessment: revised cross-informant syndromes and new DSM-oriented scales for the CBCL, YSR, and TRF: comment on lengua, sadowski, friedrich, and fischer (2001). *J Consult Clin Psychol* (2001) **69**(4):699–702. doi:10.1037/0022-006X.69.4.699
 105. Rich EC, Loo SK, Yang M, Dang J, Smalley SL. Social functioning difficulties in ADHD: association with PDD risk. *Clin Child Psychol Psychiatry* (2009) **14**(3):329–44. doi:10.1177/1359104508100890
 106. Conners CK, Epstein JN, Angold A, Klaric J. Continuous performance test performance in a normative epidemiological sample. *J Abnorm Child Psychol* (2003) **31**(5):555–62. doi:10.1023/A:1025457300409
 107. Glahn DC, Therman S, Manninen M, Huttunen M, Kaprio J, Lonnqvist J, et al. Spatial working memory as an endophenotype for schizophrenia. *Biol Psychiatry* (2003) **53**(7):624–6. doi:10.1016/S0006-3223(02)01641-4
 108. Ivry RB, Robertson LC. *The Two Sides of Perception*. Cambridge, MA: MIT Press (1998).
 109. Zaidel E, editor. Language, dichotic listening and the disconnected hemispheres. *Conference on Human Brain Function*. Los Angeles, CA: BRI Publications Office, UCLA (1976).
 110. Zaidel E. Disconnection syndrome as a model for laterality effects in the normal brain. In: Hellige J, editor. *Cerebral Hemisphere Asymmetry: Method, Theory, and Application*. New York, NY: Praeger (1983). p. 95–151.
 111. Grimshaw GM, Kwasny KM, Covell E, Johnson RA. The dynamic nature of language lateralization: effects of lexical and prosodic factors. *Neuropsychologia* (2003) **41**:1008–19. doi:10.1016/S0028-3932(02)00315-9
 112. Bryden MP, MacRae L. Dichotic laterality effects obtained with emotional words. *Neuropsychiatry Neuropsychol Behav Neurol* (1988) **1**(3):171–6.
 113. Milner AD, Goodale MA. *The Visual Brain in Action*. New York, NY: Oxford University Press (1995). 248 p.
 114. Milner AD, Goodale MA. Two visual systems re-viewed. *Neuropsychologia* (2008) **46**(3):774–85. doi:10.1016/j.neuropsychologia.2007.10.005
 115. Singh-Curry V, Husain M. The functional role of the inferior parietal lobe in the dorsal and ventral stream dichotomy. *Neuropsychologia* (2009) **47**(6):1434–48. doi:10.1016/j.neuropsychologia.2008.11.033
 116. Segal E, Petrides M. The anterior superior parietal lobule and its interactions with language and motor areas during writing. *Eur J Neurosci* (2012) **35**(2):309–22. doi:10.1111/j.1460-9568.2011.07937.x
 117. Pazzaglia M, Smania N, Corato E, Aglioti SM. Neural underpinnings of gesture discrimination in patients with limb apraxia. *J Neurosci* (2008) **28**(12):3030–41. doi:10.1523/JNEUROSCI.5748-07.2008
 118. Mort DJ, Malhotra P, Mannan SK, Rorden C, Pambakian A, Kennard C, et al. The anatomy of visual neglect. *Brain* (2003) **126**(Pt 9):1986–97. doi:10.1093/brain/awg200
 119. Geng JJ, Vossel S. Re-evaluating the role of TPJ in attentional control: contextual updating? *Neurosci Biobehav Rev* (2013) **37**(10 Pt 2):2608–20. doi:10.1016/j.neubiorev.2013.08.010
 120. Carter RM, Huettel SA. A nexus model of the temporal-parietal junction. *Trends Cogn Sci* (2013) **17**(7):328–36. doi:10.1016/j.tics.2013.05.007
 121. Klimesch W, Sauseng P, Hanslmayr S. EEG alpha oscillations: the inhibition-timing hypothesis. *Brain Res Rev* (2007) **53**(1):63–88. doi:10.1016/j.brainresrev.2006.06.003
 122. Palva S, Palva JM. New vistas for alpha-frequency band oscillations. *Trends Neurosci* (2007) **30**(4):150–8. doi:10.1016/j.tins.2007.02.001
 123. Shaw JC. Intention as a component of the alpha-rhythm response to mental activity. *Int J Psychophysiol* (1996) **24**(1–2):7–23. doi:10.1016/S0167-8760(96)00052-9
 124. Bekisz M, Wrobel A. Attention-dependent coupling between beta activities recorded in the cat's thalamic and cortical representations of the central visual field. *Eur J Neurosci* (2003) **17**(2):421–6. doi:10.1046/j.1460-9568.2003.02454.x
 125. Deiber MP, Missonnier P, Bertrand O, Gold G, Fazio-Costa L, Ibanez V, et al. Distinction between perceptual and attentional processing in working memory tasks: a study of phase-locked and induced oscillatory brain dynamics. *J Cogn Neurosci* (2007) **19**(1):158–72. doi:10.1162/jocn.2007.19.1.158
 126. Wrobel A. Beta activity: a carrier for visual attention. *Acta Neurobiol Exp* (2000) **60**(2):247–60.
 127. Bryden MP, MacRae L. Dichotic laterality effects obtained with emotional words. *Neuropsychiatry Neuropsychol Behav Neurol* (1989) **1**(3):171–6.
 128. Wang E, Ding YC, Flodman P, Kidd JR, Kidd KK, Grady DL, et al. The genetic architecture of selection at the human dopamine receptor D4 (DRD4) gene locus. *Am J Hum Genet* (2004) **74**(5):931–44. doi:10.1086/420854
 129. Ray WJ, Cole HW. EEG alpha activity reflects attentional demands, and beta activity reflects emotional and cognitive processes. *Science* (1985) **228**(4700):750–2. doi:10.1126/science.3992243
 130. Schutter DJ, Putman P, Hermans E, van Honk J. Parietal electroencephalogram beta asymmetry and selective attention to angry facial expressions in healthy human subjects. *Neurosci Lett* (2001) **314**(1–2):13–6. doi:10.1016/S0304-3940(01)02246-7
 131. Amunts K, Armstrong E, Malikovic A, Homke L, Mohlberg H, Schleicher A, et al. Gender-specific left-right asymmetries in human visual cortex. *J Neurosci* (2007) **27**(6):1356–64. doi:10.1523/JNEUROSCI.4753-06.2007
 132. McGivern RF, Adams B, Handa RJ, Pineda JA. Men and women exhibit a differential bias for processing movement versus objects. *PLoS One* (2012) **7**(3):e32238. doi:10.1371/journal.pone.0032238
 133. Abramov I, Gordon J, Feldman O, Chavarga A. Sex & vision I: spatio-temporal resolution. *Biol Sex Differ* (2012) **3**(1):20. doi:10.1186/2042-6410-3-20
 134. McGivern RF, Mutter KL, Anderson J, Wideman G, Bodnar M, Huston PJ. Gender differences in incidental learning and visual recognition memory: support for a sex difference in unconscious environmental awareness. *Pers Indiv Differ* (1998) **25**(2):223–32. doi:10.1016/S0191-8869(98)00017-8
 135. Rotenberg VS. The peculiarity of the right-hemisphere function in depression: solving the paradoxes. *Prog Neuropsychopharmacol Biol Psychiatry* (2004) **28**(1):1. doi:10.1016/S0278-5846(03)00163-5
 136. Cloninger CR, Svrakic DM, Przybeck TR. A psychobiological model of temperament and character. *Arch Gen Psychiatry* (1993) **50**(12):975–90. doi:10.1001/archpsyc.1993.01820240059008
 137. Mesulam M-M. Large-scale neurocognitive networks and distributed processing for attention, language, and memory. *Ann Neurol* (1990) **28**:597–613. doi:10.1002/ana.410280502
 138. Grefkes C, Wang LE, Eickhoff SB, Fink GR. Noradrenergic modulation of cortical networks engaged in visuomotor processing. *Cereb Cortex* (2010) **20**(4):783–97. doi:10.1093/cercor/bhp144
 139. Simos PG, Fletcher JM, Sarkari S, Billingsley RL, Denton C, Papanicolaou AC. Altering the brain circuits for reading through intervention: a magnetic source imaging study. *Neuropsychology* (2007) **21**(4):485–96. doi:10.1037/0894-4105.21.4.485
 140. Rucklidge JJ. Gender differences in attention-deficit/hyperactivity disorder. *Psychiatr Clin North Am* (2010) **33**(2):357–73. doi:10.1016/j.psc.2010.01.006
 141. Notarnicola A, Maccagnano G, Pesce V, Tafuri S, Novielli G, Moretti B. Visual-spatial capacity: gender and sport differences in young volleyball and tennis athletes and non-athletes. *BMC Res Notes* (2014) **7**:57. doi:10.1186/1756-0500-7-57

142. Shaywitz SE, Shaywitz BA. Paying attention to reading: the neurobiology of reading and dyslexia. *Dev Psychopathol* (2008) **20**(4):1329–49. doi:10.1017/S0954579408000631
143. Goldberg E, Podell K, Lovell M. Lateralization of frontal lobe functions and cognitive novelty. *J Neuropsychiatry Clin Neurosci* (1994) **6**(4):371–8. doi:10.1176/jnp.6.4.371
144. Drummond SP, Meloy MJ, Yanagi MA, Orff HJ, Brown GG. Compensatory recruitment after sleep deprivation and the relationship with performance. *Psychiatry Res* (2005) **140**(3):211–23. doi:10.1016/j.psychres.2005.06.007
145. Mercer Moss FJ, Baddeley R, Canagarajah N. Eye movements to natural images as a function of sex and personality. *PLoS One* (2012) **7**(11):e47870. doi:10.1371/journal.pone.0047870
146. Cho SC, Hwang JW, Lyoo IK, Yoo HJ, Kin BN, Kim JW. Patterns of temperament and character in a clinical sample of Korean children with attention-deficit hyperactivity disorder. *Psychiatry Clin Neurosci* (2008) **62**(2):160–6. doi:10.1111/j.1440-1819.2008.01749.x
147. Lynn DE, Lubke G, Yang M, McCracken JT, McGough JJ, Ishii J, et al. Temperament and character profiles and the dopamine D4 receptor gene in ADHD. *Am J Psychiatry* (2005) **162**(5):906–13. doi:10.1176/appi.ajp.162.5.906
148. Bussing R, Fernandez M, Harwood M, Wei H, Garvan CW, Eyberg SM, et al. Parent and teacher SNAP-IV ratings of attention deficit hyperactivity disorder symptoms: psychometric properties and normative ratings from a school district sample. *Assessment* (2008) **15**(3):317–28. doi:10.1177/1073191107313888
149. Wechsler D. *Wechsler Intelligence Scale – Revised*. San Antonio, TX: Psychological Corporation (1981).
150. Dunn LM, Markwardt FC. *Examiner's Manual: Peabody Individual Achievement Test*. Circle Pines, MN: American Guidance Service (1970).
151. Brown MB, Giandemoto MJ, Bolen LM. Diagnosing written language disabilities using the Woodcock-Johnson tests of educational achievement-revised and the Wechsler individual achievement test. *Psychol Rep* (2000) **87**(1):197–204. doi:10.2466/PRO.87.5.197-204
152. Stroop JR. Studies of interference in serial verbal reactions. *J Exp Psychol Gen* (1935) **18**:643–62. doi:10.1037/h0054651
153. Golden C. *Stroop Color and Word Test: A Manual for Clinical and Experimental Uses*. Chicago IL: Skoelting (1978).
154. Crowe SF. The differential contribution of mental tracking, cognitive flexibility, visual search, and motor speed to performance on parts A and B of the trail making test. *J Clin Psychol* (1998) **54**(5):585–91. doi:10.1002/(SICI)1097-4679(199808)54:5<585::AID-JCLP4>3.3.CO;2-M

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2015 Hale, Wiley, Smalley, Tung, Kaminsky, McGough, Jaini and Loo. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.